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(54) Title: SERINE PROTEASE INHIBITORS

(57) Abstract

The invention relates to a compound having the formula (I): R¹SO₂-B-X-Z-C(O)-Y, B is a bond, an I amino acid of the formula -NH-CH[(CH₂)_pC(O)OH]-C(O)— or an ester derivative thereof wherein p is 1, 2, or 3, Gly, D-1-Piq, D-3-Piq, D-1-Tiq, D-3-Tiq, D-Atc, Aic, or a L- or D-amino acid having a hydrophobic, basic or neutral side chain; X is an amino acid with a hydrophobic side chain, glutamine, serine, threonine, a cyclic amino acid optionally containing an additional heteroatom selected from N, O or S, and optionally substituted with (1-6C)alkyl, (1-6C)alkoxy, benzyloxy or oxo, or X is 2-amino-isobutyric acid, -NR²-CH₂-CC(O)— or the fragment (I) or (II), wherein n is 2, 3, or 4, W is CH or N and R³ is H, (1-6C)alkyl or phenyl which groups may optionally be substituted with hydroxy, (1-6C)alkoxy, COOH, COO(1-6C)alkyl, CONH₂, or halogen; Z is lysine or 4-aminocyclohexylglycine. The ecompounds of the invention have anticoagulant activity and can be used in treating or preventing thrombin-related diseases. The variable R¹¹ and Y are defined in claim

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SERINE PROTEASE INHIBITORS

The invention relates to new serine protease inhibitors, pharmaceutical compositions containing the same, as well as to the use of said inhibitors for the manufacture of a medicament for treating and preventing thrombin-related diseases.

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Serine proteases are enzymes which, amongst other things, play an importantit role in the blood coagulation cascade. Members of this group of proteases are for example thrombin, trypsin, factors VIIa, IXa, Xa, XIa, XIIa, and protein C.

Thrombin is the serine protease which regulates the last step in the coagulattion cascade. The prime function of thrombin is the cleavage of fibrinogen to generate fibrin 1 monomers, which form an insoluble gel by cross-linking. In addition, thrombin regulates its own production by activating factors V and VIII earlier in the cascade. It also has important actions at the cellular level, where it acts on specific receptors to cause platelet aggregation, endothelial cell activation and fibroblast proliferation. Thus thrombin has a central reegulatory role in haemostasis and thrombus formation. Since inhibitors of thrombin may havee a wide range of therapeutical applications, extensive research has been performed in this area.

In the development of synthetic inhibitors of serine proteases, and moore specifically of thrombin, the interest in small synthetic peptides that are recognized by protecolytic enzymes in a manner similar to that of natural substrates, has increased. As a result, new peptide-like inhibitors have been prepared, such as the transition state inhibitors of thrombin.

The search for more effective and more selective thrombin inhibitors contitinues unabated in order to obtain thrombin inhibitors which can be administered in lower dosagges and which have fewer and less severe side effects. Furthermore, special attention is paid to coral bioavailability. Potent intravenous thrombin inhibitors are clinically effective in acute care settings requiring the treatment of thrombin-related diseases. However, particularly the prevention of thrombin-related diseases such as myocardial infarct; thrombosis and stroke require ldong-term therapy, preferably by orally dosing an anticoagulant.

Many of the peptide-like serine protease inhibitors, in particular thrombin inhibitors, disclosed in prior publications are based on the sequence -D-Phe-Pro-Arg-, see for example compounds as disclosed by Brady et al. [Bioorganic & Medical Chemistry, 3 (1995), 10063-78] and in US Patent 5,597,804. Thrombin inhibitors may also contain lysine side chains innstead of arginine,

such as other inhibitors disclosed by Brady et al., and Lewis et al. [[Thrombosis and Haemostasis, 74(4) (1995), 1107-12], and further by Jones et al. [J. Enzyyme Inhibition, 9 (1995), 43-60]. In the latter publication it was reported that tripeptide compounds containing α-keto methyl ester functions are labile compounds and therefore unfavouurable for further development as thrombin inhibitors. Also thrombin inhibitors having an aminoocyclohexyl moiety instead of lysine or arginine side chain are known [WO 94/25051]. From theese and also other references [e.g. US Patent 5,523,308] a number of variations at the C-tterminus of these peptide-like thrombin inhibitors is known. The developments in this field have: already improved the understanding of how to modulate the biological properties of this ttype of thrombin inhibitors. However, although great effort is being spend on finding seelective thrombin inhibitors having good oral bioavailability there are still few transition state thhrombin inhibitors known in the art which fulfill these requirements.

Surprisingly, it has now been found that compounds of the formula:

 $R^{1}SO_{2}-B-X-Z-C(O)-Y$ (I)

wherein R¹ is R²OOC-(CHR²)_m- or R²NH-CO-(CHR²)_m- or is selected from (1-12C)alkyl, (2-12C)alkenyl, which groups may optionally be substituted with ([3-12C)cycloalkyl, (1-6C)alkoxy, OH, COOR², CF₃ or halogen, and from (6-14C)aryl, (77-15C)aralkyl and (8-16)aralkenyl, the aryl groups of which may optionally be substituted with (1-6C)alkyl, (3-8C)cycloalkyl, (1-6C)alkoxy, OH, COOH, CF₃ or halogen;

m is 1, 2 or 3;

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WO 98/50420

each group R² is independently H, (1-12C)alkyl, (3-8C)cycloalkyl, (6-14C)aryl or (7-15C)aralkyl, the aryl groups of which may be substituted with (1-6C)alkyl, (1-6C)alkoxy or halogen;

B is a bond, an amino-acid of the formula -NH-CH[(CH₂)_pC(O)OH]-Cl(O)- or an ester derivative thereof wherein p is 1, 2 or 3, Gly, D-1-Piq, D-3-Piq, D-1-Tiq, D-33-Tiq, D-Atc, Aic, or a L- or D-amino acid having a hydrophobic, basic or neutral side chain;

X is an amino acid with a hydrophobic side chain, glutamine, serine, threonine, a cyclic amino acid optionally containing an additional heteroatom selected from N, O or S, and optionally substituted with (1-6C)alkyl, (1-6C)alkoxy, benzyloxy or oxo, or X is 2-aminno-isobutyric acid, -NR²-CH₂-C(O)- or the fragment

-NH-CH Or Or
$$\mathbb{R}^3$$

W \mathbb{R}^3
-NH-CH₂-C(O)-
or \mathbb{R}^3

wherein n is 2, 3, or 4, W is CH or N and R³ is H, (1-6C)alkyl or phenyl which groups may optionally be substituted with hydroxy, (1-6C)alkoxy, COOH, COO(1-6C)allkyl, CONH₂, or halogen;

5 Z is lysine or 4-aminocyclohexylglycine;

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Y is -NH-(1-6C)alkylene- C_6H_5 , the phenyl group of which may be substituted with (1-6C)alkyl, (1-6C)alkoxy or halogen, or Y is -OR⁴ or -NR⁵R⁶, wherein R⁴ is H, (2-6C)alkyl or benzyl, and R⁵ and R⁶ are independently H, (1-6C)alkoxy, or (1-6C)alkyl optionally substituted with halogen, or R⁵ and R⁶ together are (3-6C)alkylene, or R⁵ and R⁶ together with the nitrogen

atom to which they are bonded are view wherein V is O, S or SO₂; or a prodrug thereof or a pharmaceutically acceptable salt thereof,

are potent and selective serine protease inhibitors. Specifically, the compounds of the present invention are inhibitors of thrombin, of factor VIIa/tissue factor and of factor \cdot Xa. Compounds of the invention show improved pharmacokinetics, and in particular good bicoavailability after oral administration. The α -(2-6C)keto ester compounds which are part of the present invention do not show the disadvantages of the previously reported labile α -keeto methyl ester compounds.

The compounds of the present invention are useful for treating and preventing thrombin-mediated and thrombin-associated diseases. This includes a number of thrombotic and prothrombotic states in which the coagulation cascade is activated which include, but are not limited to, deep vein thrombosis, pulmonary embolism, thrombophlebitis, aarterial occlusion from thrombosis or embolism, arterial reocclusion during or after angioplasty, or thrombolysis, restenosis following arterial injury or invasive cardiological procedures, posttoperative venous thrombosis or embolism, acute or chronic atherosclerosis, stroke, myocardial i infarction, cancer and metastasis, and neurodegenerative diseases. The compounds of the invention may also be used as anticoagulants in extracorporeal blood circuits, as necessary in dialysis: and surgery.

The compounds of the invention may also be used as in vitro anticoagulants.

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Preferred serine protease inhibitors according to this invention are the compounds wherein Z is lysine. More preferred are the compounds wherein X is a cyclic amino acid, an I amino acid with a hydrophobic side chain, glutamine, serine, threonine, -NR²-CH₂-C(O)-, or the fragment

$$R^3$$
-NH-CH N-CH₂-C(O)-
Or Or O

wherein R³ is H, (1-6C)alkyl or phenyl.

Particularly preferred are the compounds wherein X is proline, leucine, glutaamine, threonine, phenylalanine, -NR²-CH₂-C(O)- wherein R² is methyl, cyclopentyl or cycclohexyl, or the fragment

$$R^3$$
-NH-CH N-CH₂-C(O)-
O
O
O
 N -CH₂-C((O)-

wherein R³ is H or methyl.

Other preferred compounds are those wherein B is a bond or a D-amingo acid having a hydrophobic or neutral side chain. The most preferred compounds of the invention are those wherein R¹ is (1-6C)alkyl or benzyl. Preferably R⁴ in the definition of Y is (2-6C)alkyl or benzyl. In particular preferred are the compounds wherein Y is -OCH(CH₃)₂. Also preferred compounds have Y is NH₂.

The term (1-12C)alkyl means a branched or unbranched alkyl group havingg 1 to 12 carbon atoms, such as methyl, ethyl, t-butyl, isopentyl, heptyl, dodecyl, and the likee. Preferred alkyl groups are (1-6C)alkyl groups, having 1-6 carbon atoms.

A (2-12C)alkenyl group is a branched or unbranched unsaturated hydrocarbonn group having 2 to 12 carbon atoms. Preferred are (2-6C)alkenyl groups. Examples are ethenyyl, propenyl, allyl, and the like.

The term (1-6C)alkylene means a branched or unbranched alkylene group having 1 to 6 carbon atoms, such as -(CH₂)_s- and s is 1 to 6, -CH(CH₃)-, -CH(CH₃)-, etc. PPreferred alkylene groups in the definition of Y are ethylene and methylene.

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The term (1-6C)alkoxy means an alkoxy group having 1-6 carbon atoms, thee alkyl moiety of which has the meaning as previously defined.

The term (3-12C)cycloalkyl means a mono- or bicycloalkyl group having 3-112 carbon atoms which cycloalkyl group may optionally be substituted with an oxo groupp. Preferred are (3-8C)cycloalkyl, such as cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cycloheptyl, cyclopentyl, etc.. Cyclopentyl and cyclohexyl are even more preferred cycloalkyl groups. A preferred cycloalkyl substituted alkyl group in the definition of R¹ is the camphor group.

A (6-14C)aryl group is an aromatic moiety of 6 to 14 carbon atoms. The aryl ggroup may further contain one or more hetero atoms, such as N, S, or O. Examples of aryl groups are phenyl, naphthyl, (iso)quinolyl, indanyl, and the like.

(7-15C)Aralkyl and (8-16C)aralkenyl groups are alkyl and alkenyl groups respectively, substituted by one or more aryl groups, the total number of carbon atoms beining 7 to 15 and 8 to 16, respectively.

The term halogen means fluorine, chlorine, bromine or iodine.

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The term ester derivative means any appropriate ester derivative, preferably (11-4C)alkyl-esters, such as methyl-, ethyl- or t-butyl-esters.

The terms Atc means 2-aminotetralin-2-carboxylic acid and Aic means amino i indane carboxylic acid. The terms 1- and 3-Tiq mean 1,2,3,4-tetrahydroisoquinoline-1- and -33-carboxylic acid, respectively; 1- and 3-Piq are perhydroisoquinoline-1- and -3-carboxylic acid, respectively.

The term amino acid having a hydrophobic side chain means an amino acid haaving a side chain being (3-8C)cycloalkyl, (6-14C)aryl or (1-6C)alkyl, which alkyl group mnay optionally be substituted with one or more (3-8C)cycloalkyl groups or (6-14C)aryl groups. The hydrophobic side chain may optionally be substituted with one or more substituents, such as hydroxy, halogen, trifluoromethyl, -OSO₂CF₃, (1-4C)alkyl (for instance methyl or ethnyl), (1-4C)alkoxy (for instance methoxy), phenyloxy, benzyloxy, and the like. Preferred amnino acids with a hydrophobic side chain are leucine, valine, cyclohexylalanine, 4-methoxy-ccyclohexylalanine, cyclo-octylalanine, phenylalanine, D-naphthylalanine, tyrosine, O-methyl tyrosine (or: p-methoxy-phenylalanine), 3,3-diphenylalanine, norleucine and leucine.

Amino acids having a basic side chain are for example, but not limited to, anginine and lysine, preferably arginine.

The term amino acids having a neutral side chain refers to amino acids such ass glutamine (Gln), methionine sulfon, asparagine (Asn) and the like. Preferred are Gln and Asn.

Cyclic amino acids are for example 2-azetidine carboxylic acid, proline, pipecoblic acid, 1-amino-1-carboxy-(3-8C)cycloalkane (preferably 4C, 5C or 6C), 4-piperidine carboxylic acid, 4-thiazolidine carboxylic acid, 3,4-dehydro-proline, azaproline, 2-octahydrobindole carboxylic acid, and the like. Preferred are 2-azetidine carboxylic acid, proline,; pipecolic acid, 4-thiazolidine carboxylic acid, 3,4-dehydro-proline and 2-octahydroindole carbboxylic acid. In the definitions, the term substituted means: substituted by one or more substituents.

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The invention also includes prodrugs of the compounds of formula I, which affter administration are metabolized into the active compounds. Suitable prodrugs are 1 for example Nalkoxycarbonyl protected (preferably N-ethoxycarbonyl) derivatives of the compounds of formula I.

The invention further includes a process for preparing a compound of formnula I, comprising coupling of suitably protected amino acids or amino acid analogs, followed! by removing the protective groups.

The compounds according to formula I may be prepared in a manner convventional for such compounds. To that end, suitably Na protected (and side-chain protected if reeactive side-chains are present) amino acid derivatives or peptides are activated and coupled to suitably carboxyl protected amino acid or peptide derivatives either in solution or on a solid suupport. Protection of the α -amino functions generally takes place by urethane functions such as the acid-labile tertbutyloxycarbonyl group (Boc), benzyloxycarbonyl (Cbz) group and substitutited analogs or the base-labile 9-fluorenyl-methyloxycarbonyl (Fmoc) group. The Cbz group cann also be removed by catalytic hydrogenation. Other suitable amino protective groups include: Nps, Bpoc, Msc, etc. A good overview of amino protective groups is given is given in The PPeptides, Analysis, Synthesis, Biology, Vol. 3 E. Gross and J. Meienhofer, Eds., (Academic FPress, New York, 1981). Protection of carboxyl groups can take place by ester formation e.g.; base-labile esters like methyl- or ethylesters, acid labile esters like tert-butylesters, or hydrogenolytically-labile esters like benzylesters. Protection of the side chain function oof lysine or 4aminocyclohexylglycine may be accomplished by using the aforementioned ¿groups. Activation of the carboxyl group of the suitably protected amino acids or peptides cann take place by the azide, mixed anhydride, active ester, or carbodiimide method, especially with the addition of catalytic and racemization-suppressing compounds like 1-hydroxybbenzotriazole, 3-hydroxy-4-oxo-3,4-dihydro-1,2,3-benzotriazine, N-hydroxy-5-norhydroxysuccinimide,

bornene-2,3-dicarboximide. See, e.g. The Peptides, Analysis, Synthesis, Bioblogy (see above) and Pure and Applied Chem. 59(3), 331-344 (1987).

The compounds of the invention, which can be in the form of a free base, mayy be isolated from the reaction mixture in the form of a pharmaceutically acceptable salt. The: pharmaceutically acceptable salts may also be obtained by treating the free base of formula I with an organic or inorganic acid such as hydrogen chloride, hydrogen bromide, hydrogen iodilide, sulfuric acid, phosphoric acid, acetic acid, propionic acid, glycolic acid, maleic acid, malonic acid, methanesulfonic acid, fumaric acid, succinic acid, tartaric acid, citric acid, bbenzoic acid, and ascorbic acid.

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The compounds of this invention possess one or more chiral carbon atoms, and may therefore be obtained as a pure enantiomer, or as a mixture of enantiomers, or as a mixture containing diastereomers. Methods for obtaining the pure enantiomers are well known in the art, e.g. crystallization of salts which are obtained from optically active acids and the racemic mixture, or chromatography using chiral columns. For diastereomers straight phase our reversed phase columns may be used.

The compounds of the invention may be administered enterally or parenterallyy, and for humans preferably in a daily dosage of 0.001-100 mg per kg body weight, preferably 00.01-10 mg per kg body weight. Mixed with pharmaceutically suitable auxiliaries, e.g. as described in the standard reference, Gennaro et al., Remington's Pharmaceutical Sciences, (18th ed., Mack Publishing Company, 1990, see especially Part 8: Pharmaceutical Preparations and Their: Manufacture) the compounds may be compressed into solid dosage units, such as pills, tabletss, or be processed into capsules or suppositories. By means of pharmaceutically suitable liquidds the compounds can also be applied in the form of a solution, suspension, emulsion, e.g. for use as an injection preparation, or as a spray, e.g. for use as a nasal spray.

For making dosage units, e.g. tablets, the use of conventional additives such as fillers, colorants, polymeric binders and the like is contemplated. In general any pharmaceunically acceptable additive which does not interfere with the function of the active compounds caan be used.

Suitable carriers with which the compositions can be administered includde lactose, starch, cellulose derivatives and the like, or mixtures thereof, used in suitable amounts.

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The invention is further explained by reference to the following illustrative Examples.

GENERAL

Abbreviations:

5 Et = ethyl Bzl = benzyl

Boc = tert-butyloxycarbonyl Cbz = benzyloxycarbonyl

Cha = cyclohexylalanyl Pro = prolyl

Lys = lysyl Acg = 4-aminocyclohexyl glycyl

TFA = trifluoro acetic acid Pac = phenylacetyl

10 Nps = nitrophenylsulfonyl Bpoc = 2-p-biphenylisopropyloxycarboonyl

Asp = aspartyl Glu = glutamyl

Dpa = diphenylalanyl H-Aad-OH = amino-adipic acid

Tyr(Me) = (O-methyl)-tyrosyl Phe = phenylalanyl

Nal = naphthylen-2-yl-alaninyl

15 H-3-Tiq-OH = 1,2,3,4-tetrahydroisoquinoline-3-carboxylic acid

Msc = methylsulfonylethyloxycarbonyl

Teoc = 2-(trimethylsilyl)ethoxycarbonyl

norLeu(cyclo)-Gly-OH = (S)-3-amino-2-oxo-hexahydro-1-azepineacetic acid

norVal(cyclo)-Gly-OH = (S)-3-amino-2-oxo-1-piperidineacetic acid

Experimental:

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The solvent systems used in HPLC are:

A: 0.5 M phosphate buffer pH = 2.1; B: water; C: acetonitrile/water 3/2 v/v.

Unless stated otherwise the retention times (Rt (LC)) were determined on ann analytical HPLC Supelcosil LC-18-DB column (5 µm particles; 250 x 2.1 mm), which was eluted using a gradient (as specified) of solvent systems A, B and C at a flow rate of 0.25 ml//min at 35 °C.

Example 1.

BzISO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-NHBzl

30 (a) Cbz-Lys(Boc)-OMe

To a solution of Cbz-Lys(Boc)-OH (28 g) in dichloromethane/methanol (9/1 v/v; 500 mL) was added 2-(1H-benzotriazol-1-yl)-1,1,3,3-tetramethyluronium tetrafluoroboratee (23.6 g) and the solution was adjusted to pH 8 by addition of triethylamine. The reaction mixture was stirred for

2 hours at room temperature. The mixture was washed successively with cold 1 IN hydrochloric acid, water, 5% sodium hydrogencarbonate, and water and dried over sodilium sulfate. The filtrate was evaporated and the residue was chromatographed on silica gel using ethyl acetate/heptane (1/4 v/v) as eluent. The fractions containing Cbz-Lys(Boc)-O)Me were pooled and evaporated. Yield: 29.1 g

TLC: Rf= 0.85, ethyl acetate/heptane=3/1 v/v on silica.

(b) Cbz-Lys (Boc)Ψ[cyanoacetate]

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To a cold (-78°C) solution of Cbz-Lys(Boc)-OMe (29.1 g) in dry dichloromethane (800 mL) was added dropwise diisobutylaluminium hydride (222 mL of 1M solution in 1 hexane) keeping the reaction temperature below -70° C. The resulting solution was stirred at --78° C for 1 hour and an aqueous 5% citric-acid solution (600 mL) was added to the reaction rmixture. The two layer mixture was stirred at room temperature for 10 minutes, the layers were separated and the aqueous layer was extracted twice with dichloromethane. The combined dichloromethane layers were washed with water, dried over sodium sulfate and filtered. The filtrate wwas stirred under a nitrogen atmosphere and cooled on a icewater-bath. A solution of sodium cyaanide (36.3 g) and benzyltriethyl ammonium chloride (4.2 g) in water (600 mL) was added. Underr vigorous stirring acetic anhydride was added portionwise (2 x 9 mL) over a period of 30 min. The organic layer was separated and the aqueous layer was extracted twice with dichloromethane. The combined dichloromethane layers were washed with water, dried over sodium sulfifate, filtered and evaporated in vacuo. The residue was purified by chromatography onn silica (eluent : heptane/ethyl acetate= 1/1 v/v) to yield Cbz-Lys (Boc) \P[cyanoacetate] (26.3 ; g.)

TLC: Rf = 0.60, dichloromethane/ethyl acetate = 7/3 v/v on silica.

25 (c) Cbz-Lys(Boc)Ψ[CHOHCO]-OMe

A solution of Cbz-Lys(Boc)Ψ[cyanoacetate] (26.3 g.) in diethylether/methanool = 3/1 v/v (600 mL) was cooled to -20° C under a nitrogen atmosphere, and 66 g of gaseous I hydrogen chloride was introduced keeping the temperature below -5° C. The reaction mixture: was kept at 4° C overnight. Water (100 mL) was added dropwise to the reaction mixture keeping the temperature below 5° C. After stirring for 16 h at room temperature the corganic layer was separated and washed with water. The aqueous layer was saturated with soodium chloride and extracted with sec-butanol/dichloromethane = 3/2 v/v. The organic phase was washed with brine, dried over sodium sulfate, filtered and evaporated in vacuo to give 2.5.4 g of the crude

amine. The residue was taken up in N,N-dimethylformamide (400 mL), di-tert-l-butyl dicarbonate (16 g) was added and adjusted to pH 8 using triethylamine. The reaction mixture was stirred at room temperature overnight. The solvent was removed by evaporation at reducced pressure. The residue was dissolved in ethyl acetate, washed with water and brine successsively, dried over sodium sulfate, filtered and evaporated in vacuo. The residue was purified byy chromatography on silica (eluent: ethyl acetate/heptane = 4/6 v/v) to yield Cbz-Lys(Boc)Y[[CHOHCO]-OMe (15.8 g).

TLC: Rf = 0.75, ethyl acetate/pyridine/acetic acid/water=63/20/6/11 v/v/v/v onn silica.

10 (d) Cbz-Lys(Boc)Ψ[CHOHCO]-OH

A stirred solution of Cbz-Lys(Boc)Y[CHOHCO]-OMe (2.0 g) in dioxane/wvater=7/3 v/v (50 mL) at room temperature was treated portionwise with a 2M sodium hydroxide solution (2.36 mL). After 1 hour the reaction mixture was diluted with water (100 mL), 2M I hydrochloric acid was added until pH 2.0 and extracted with dichloromethane. The combined organic phases were washed with water, dried over sodium sulfate, filtered and concentrated in vaacuo to yield Cbz-Lys(Boc)Y[CHOHCO]-OH (1.85 g).

TLC: Rf=0.65, ethyl acetate/pyridine/acetic acid/water=63/20/6/11 v/v/v/v on silica.

(e) Cbz-Lys(Boc)Ψ[CHOHCO]-NHBzl

To a stirred solution of Cbz-Lys(Boc)Ψ[CHOHCO]-OH (0.90 g) in N,N-dilimethylformamide (10 mL) were added 1-hydroxybenzotriazole (HOBt, 444 mg), N-methylmorpholine (0.5 mL), benzylamine (282 mg) and 1-(3-dimethylaminopropyl)-3-ethylcarbodiimidde hydrochloride (EDCI, 462 mg). After stirring for 16 hours at room temperature the reacction mixture was poured into water and this aqueous mixture was extracted with ethyl acetate.. The ethyl acetate extract was washed with 1N hydrochloric acid, water, aqueous 5% sodium hhydrogencarbonate and water, dried over sodium sulfate, filtered and concentrated in vacuuo to yield Cbz-Lys(Boc)Ψ[CHOHCO]-NHBzl (1.0 g).

TLC: Rf=0.81, ethyl acetate/pyridine/acetic acid/water=163/20/6/11 v/v/v/v opn silica.

30 (f) H-Lys(Boc)Ψ[CHOHCO]-NHBzl.HCl

To a solution of Cbz-Lys(Boc)Ψ[CHOHCO]-NHBzl (1.0 g) in methanol (2.5 mL) were added 10% palladium on activated carbon (100 mg) and 2M hydrochloric acidi (1 mL) and this

suspension was hydrogenated at atmospheric pressure for 1 hour at room temperature. The palladium catalyst was removed by filtration and the filtrate was concentrated I in vacuo to yield H-Lys(Boc) Y[CHOHCO]-NHBzl. HCl (0.87 g).

TLC: Rf=0.15, ethyl acetate/pyridine/acetic acid/water=163/20/6/11 v/v/v/v onn silica.

(g) N-Boc-L-α-Amino-ε-caprolactam

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To a stirred solution of L-α-Amino-ε-caprolactam (10g) in dioxane/water (ζ2/1 v/v) (30 mL) was added 1N sodium hydroxide solution (7.8 mL) followed by di-t-butyl dicaarbonate (18.8 g). The mixture was stirred for 16 hours at room temperature and concentrated in vacuo. The residue was dissolved in ethyl acetate and washed with water and brine, ddried over sodium sulfate, filtered and evaporated in vacuo. The crude material was triturated bby hexane, filtered and dried in vacuo to yield N-Boc-L-α-Amino-ε-caprolactam (16 g).

TLC: Rf= 0.85, ethyl acetate/heptane 1/1 v/v on silica.

15 (h) Boc-norLeu(cyclo)-Gly-OMe.

N-Boc-L-α-Amino-ε-caprolactam (10 g) was dissolved in dichloromethane ((100 mL). At -20 °C a 1M solution of lithium bis (trimethylsilyl)amide in tetrahydrofuran/cyclobhexane 1/1 v/v (1 equiv.) was added slowly and the mixture was stirred for 30 min. Methyl bronmoacetate (4 mL) was subsequently added and the mixture was stirred for 2 hours at rooom temperature. Additional lithium bis (trimethylsilyl)amide in tetrahydrofuran/cyclohexane 1/11 v/v was added to force the reaction to completion. The mixture was diluted by dichloromethanee and washed with 0.1N hydrochloric acid, water, 5% aqueous sodium hydrogencarbonate scolution and brine, dried over sodium sulfate, filtered and evaporated in vacuo. The residuee was purified by chromatography on silica gel (eluent: heptane/ethyl acetate 6/4 v/v) to yield 12 g BocnorLeu(cyclo)-Gly-OMe.

TLC: Rf= 0.55, ethyl acetate/heptane 6/4 v/v on silica.

(i) BzlSO₂-norLeu(cyclo)-Gly-OMe.

Boc-norLeu(cyclo)-Gly-OMe (3 g) was dissolved in TFA/dichloromethane 1//1 v/v (30 mL) and stirred for 1 hour at room temperature. The reaction mixture was concentraated in vacuo. The residue was dissolved in dichloromethane (25 mL) and a solution of benazylsulfonylchloride (2.25 g) in dichloromethane (10 mL) was added slowly at 0 °C. Triethylammine was added to

keep the pH at 8 during the reaction. The mixture was stirred for 1 hour at rroom temperature, whereafter the mixture was concentrated in vacuo. The residue was dissolveed in ethyl acetate and washed with 5% sodium hydrogencarbonate solution, water and brine, ddried over sodium sulfate, filtered and evaporated in vacuo. The residue was purified by chromattography on silica gel (eluent: dichloromethane/ethyl acetate 95/5 v/v) to yield BzlSO₂-norLeuu(cyclo)-Gly-OMe (3.9 g)

TLC: Rf= 0.40, dichloromethane/ethyl acetate 9/1 v/v on silica.

(j) BzlSO2-norLeu(cyclo)-Gly-OH.

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A solution of BzlSO₂-norLeu(cyclo)-Gly-OMe (3.9 g) in dioxane /water 9/1 ((100 mL) at room temperature was treated with sufficient 1N sodium hydroxide to keep the pH. at 13 for 2 hours. After acidification, the mixture was poured into water and extracted with dichhloromethane. The organic layer was washed with water and dried on sodium sulfate The filtrate; was concentrated to yield 3.6 g of the title compound.

TLC: Rf= 0.60, ethyl acetate/pyridine/acetic acid/water 63/20/6/11 v/v/v/v on silica.

(k) BzlSO₂-norLeu(cyclo)-Gly-Lys(Boc)Ψ[CHOHCO]-NHBzl

To a cold (0°C) solution of BzlSO₂-norLeu(cyclo)-Gly-OH (340 I mg) in N,N-dimethylformamide (10 mL) were successively added 1-hydroxybenzotriazolee (HOBt, 203 mg) and dicyclohexylcarbodiimide (DCC, 217 mg). After stiring for 30 mininutes at 0°C H-Lys(Boc)Ψ[CHOHCO]-NHBzl. HCl (402 mg), prepared as describedd under (f), and triethylamine (0.15 mL) were added. The mixture was stirred at 0°C for 1 hour and then kept at room temperature overnight. The mixture was cooled to -20°C and dicyyclohexylurea was removed by filtration. The filtrate was evaporated to dryness. The residue wass dissolved in ethyl acetate and washed successively with 1M hydrochloric acid, water, aquueous 5% sodium hydrogencarbonate, water and brine, dried over sodium sulfate and concentrated in vacuo to afford BzlSO₂-norLeu(cyclo)-Gly-Lys(Boc)Ψ[CHOHCO]-NHBzl (690 mg).

TLC: Rf=0.75, ethyl acetate/pyridine/acetic acid/water=163/20/6/11 v/v/v/v oon silica.

30 (l) <u>BzlSO₂-norLeu(cyclo)-Gly-Lys(Boc)Ψ[COCO]-NHBzl</u>

To a solution of BzlSO₂-norLeu(cyclo)-Gly-Lys(Boc)Y[CHOHCO]-NHBzld (680 mg) in dry dichloromethane (20 mL) was added 424 mg of periodinane (Dess-Marttin reagent). After stirring at room temperature for one hour, aqueous 2% sodium thiosulfate solution (20 mL) and

aqueous 5% sodium hydrogencarbonate solution (20 mL) were added and I the mixture was stirred for 30 min at room temperature. The organic layer was separated, weashed with water, dried over sodium sulfate, filtered and evaporated in vacuo to givee crude BzISO₂-norLeu(cyclo)-Gly-Lys(Boc)Ψ[COCO]-NHBzl (561 mg).

TLC: Rf=0.85, ethyl acetate/pyridine/acetic acid/water=163/20/6/11 v/v/v/v onn silica.

(m) BzlSO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-NHBzl

BzlSO₂-norLeu(cyclo)-Gly-Lys(Boc)Ψ[COCO]-NHBzl (560 mg, crude) vwas treated with trifluoroacetic acid (10 mL) and stirred for 1 hour at room temperature. Thee reaction mixture was concentrated in vacuo and the residue dissolved in water and directly charged onto a preparative HPLC DeltaPak RP-C₁₈ column, which was subsequently elutedd using a gradient elution system of 20% A/80% B to 20% A/45% B/35% C over 45 min at a flow rate of 80 mL/min. Yield: 287 mg of BzlSO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-OH.

Rt (LC): 23.8 min; 20% A/60% B/20%C to 20% A/80% C in 30 min.

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Example 2.

EthylSO₂-D-Cha-Pro-LysΨ[COCO]-OH

(a) Boc-D-Cha-Pro-OPac

To a solution of Boc-D-Cha-OH.H₂O (21.5 g) in N,N-dimethylformamide: (143 mL) at 0°C were added hydroxybenzotriazole (HOBt) (13.7 g) and dicyclohexylcarbodiinmide (DCC) (15.7 g) and stirred at 0°C for 30 minutes. H-Pro-OPac. TFA (20 g) was disolved: in 50 mL of N,N-dimethylformamide, the pH was adjusted to 8 with triethylamine and this solution was added to the reaction mixture. This was allowed to continue for 16 hours during which the temperature was increased to room temperature. The mixture was filtered, concentrated in vacuo, dissolved in ethylacetate, washed with 1N hydrochloric acid, water, 5% sodium hhydrogencarbonate solution and brine, dried over sodium sulfate, filtered and evaporated in vacuo. Yield 28 g.

TLC: Rf= 0.5, dichloromethane/methanol 95/5 v/v on silica

(b) EthylSO2-D-Cha-Pro-OPac

Boc-D-Cha-Pro-OPac (3.8 g) was dissolved in TFA/dichloromethane 1/1 vv/v (25 mL) and stirred for 30 minutes at room temperature. The reaction mixture was evaporrated in vacuo. The crude amine was dissolved in dichloromethane (50 mL) and ethanesulfonyl chloride (0.8 mL)

14

was added at -78°C. Triethylamine was added to keep the pH at 8 during 1 the reaction. The mixture was stirred for 3 hours at 0°C, whereafter water (25 mL) was; added. After an additional stirring for 30 minutes at room temperature, the reaction mixture was concentrated in vacuo. The residue was dissolved in diethyl ether and washed with 1N hydrochhloric acid, water, 5% sodium hydrogencarbonate solution and brine, dried over sodium sullfate, filtered and evaporated in vacuo. Trituration of the crude material with methanol yielded I ethylSO₂-D-Cha-Pro-OPac (3.0 g).

TLC: Rf= 0.6, dichloromethane/methanol 95/5 v/v on silica.

10 (c) EthylSO₂-D-Cha-Pro-OH

To a solution of ethylSO₂-D-Cha-Pro-OPac (10 g) in tetrahydrofuran (250 mhL) was added 1M solution of tetrabutylammonium fluoride in tetrahydrofuran (84 mL). The reaction mixture was stirred for 30 minutes at room temperature and poured into water (1 L). Thee aqueous solution was extracted with ethyl acetate. The combined organic layers were successively washed with 1N hydrochloric acid and water, dried over sodium sulfate and concentrated in vacuo. The residue was purified by crystallisation from ethyl acetate/diisopropylether to yyield EthylSO₂-D-Cha-Pro-OH (6.0 g).

TLC: Rf= 0.2, ethyl acetate/pyridine/acetic acid/water 163/20/6/11 v/v/v/v on silica.

20 (d) EthylSO₂-D-Cha-Pro-LysΨ[COCO]-OH

The DCC/HOBt-coupling between EthylSO₂-D-Cha-Pro-OH and H-Lys(B6oc)Ψ[CHOHCO]-OMe.HCl, saponification, Dess-Martin oxidation, deprotection and purificcation were done according to the procedures described in example 1. Yield: 163 mg of the titlee compound.

Rt (LC): 36.35 min. 20% A/80% B to 20% A/20% B/60% C in 40 min.

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Example 3.

BzlSO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-OEt

(a) Cbz-Lys(Boc) Ψ[CHOHCO]-OEt

Cbz-Lys(Boc) Ψ [CHOHCO]-OMe (751 mg) was dissolved in 25 mL obf 3N HCl/ethanol solution and stirred during 4.5 hours at room temperature. The reaction solution was evaporated to dryness and coevaporated three times with ethanol to yield 6591 mg of Cbz-Lys Ψ [CHOHCO]-OEt. This product was dissolved in 10 mL dry dichloromethanne and di-tert-butyl

dicarbonate (425 mg) was added. The pH of the solution was adjusted and maaintained at 8 with triethylamine and the reaction was stirred for 16 hours at room temperature. Water was added and the organic layer was washed and dried to yield 782 mg of the desired product. After purification on silica using heptane/ethyl acetate 2/3 the final yield was 696 mgg.

5 TLC: Rf= 0.95, ethyl acetate/pyridine/acetic acid/water 232/31/18/7 v/v/v/v opn silica.

(b) H-Lys(Boc)Ψ[CHOHCO]-OEt.HCl

To a solution of Cbz-Lys(Boc)Ψ[CHOHCO]-OEt (696 mg) in ethanol (25 i mL) were added 10% palladium on activated carbon (100 mg) and 2N hydrochloric acid ((0.8 mL) and this suspension was hydrogenated at atmospheric pressure for 50 minutes at room1 temperature. The palladium catalyst was removed by filtration and the filtrate was concentrated i in vacuo to yield H-Lys(Boc)Ψ[CHOHCO]-OEt. HCl (525 mg).

TLC: Rf=0.17, ethyl acetate/pyridine/acetic acid/water=232/31/18/7 v/v/v/v onn silica.

15 (c) BziSO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-OEt

Coupling with BzlSO₂-norLeu(cyclo)-Gly-OH, oxidation, deprotection and I purification were done according to procedures described in Example 1. Yield: 186 mg of the tittle compound.

Rt (LC): 32.46 min. 20% A/80% B to 20% A/20% B/60% C in 40 min.

20 Example 4.

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BzlSO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-NH₂

The coupling between BziSO₂-norLeu(cyclo)-Gly-OH and H-Lys(Boc)\P{CHCOHCO}-NH₂.HCl. and the subsequent oxidation, deprotection and purification were done according to procedures described in Example 1 to yield 103 mg of the title compound.

25 Rt (LC): 27.50 min. 20% A/80% B to 20% A/20% B/60% C in 40 min.

Example 5.

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EthylSO₂-D-Cha-Pro-LysΨ[COCO]-OEt

The DCC/HOBt-coupling between EthylSO₂-D-Cha-Pro-OH (270 mg) and H-Lys(Boc)\(Psi(CHOHCO)\)-OEt.HCl (268 mg), Dess-Martin oxidation, deprotection using trifluoroacetic acid and purification were done according to the procedures described in Example 1. Yield: 41 mg of the title compound.

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Rt (LC): 40.7 min. 20% A/80% B to 20% A/20% B/60% C in 40 min. and maintain this mixture of eluens for an additional 10 min.

Example 6.

EthylSO₂-D-Cha-Pro-LysΨ[COCO]-NHBzl

The DCC/HOBt-coupling between EthylSO₂-D-Cha-Pro-OH (250 mg) and H-Lys(Boc)\P[CHOHCO]-NHBzl.HCl (611 mg), Dess-Martin oxidation, deeprotection using trifluoroacetic acid and purification were done according to the procedures described in Example 1. Yield: 208 mg of the title compound.

10 Rt (LC): 28.7 min. 20% A/60% B/20%C to 20% A/80% C in 30 min.

Example 7.

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EthylSO₂-D-Cha-Pro-LysΨ[COCO]-NH₂

The procedures described in Example 1 were used to prepare the titlde compound. H-Lys(Boc)Ψ[CHOHCO]-NH₂.HCl (0.84 g) was prepared from Cbz-Lys(Boc)Ψ[CHOHCO]-OH (0.95 g) as described for H-Lys(Boc)Ψ[CHOHCO]-NHBzl.HCl. Then DCCC/HOBt-coupling between EthylSO₂-D-Cha-Pro-OH (189 mg) and H-Lys(Boc)Ψ[CHOHCCO]-NH₂.HCl (179 mg), Dess-Martin oxidation, deprotection using trifluoroacetic acid and purification yielded 126 mg of the title compound.

20 Rt (LC): 36.3 min. 20% A/80% B to 20% A/20% B/60% C in 40 min.

Example 8.

BzlSO₂-norVal(cyclo)-Gly-LysΨ[COCO]-OH

(a) (S)-3-((benzyloxycarbonyl)amino)-2-oxo-piperidine

Cbz-Ornithine-OH.HCl (25 g) was dissolved in 2 L of N,N-dimethyl formamide and 12 mL of triethyl amine was added to a pH of 8.5. 2-(1H-benzotriazol-1-yl)-1,1,3,3-teetramethyluronium tetrafluoroborate (TBTU, 26.5 g) in 250 mL of N,N-dimethyl formamide was added dropwise under vigorous stirring. The mixture was allowed to react for 16 hours at room temperature while continously adjusting the pH with triethyl amine to 8.5. The reacction mixture was concentrated to dryness, dissolved in ethyl acetate and washed with 1N hydrochloric acid, water, 5% sodium hydrogen carbonate, water and brine, dried on sodium ssulfate, filtered and evaporated to dryness to yield 11.7 g of the title compound.

TLC: Rf=0.80, ethyl acetate/pyridine/acetic acid/water=63/20/6/11 v/v/v/v on ssilica.

dichloromethane/methanol 95/5 v/v) to yield 4.7 g Cbz-norVal(cyclo)-Gly-OM(e.

(b) Cbz-norVal(cyclo)-Gly-OMe.

(S)-3-((benzyloxycarbonyl)amino)-2-oxo-piperidine (5 g) was dissolved in dichhloromethane (50 mL). At -20 °C a 1M solution of lithium bis (trimethyylsilyl)amide in tetrahydrofuran/cyclohexane 1/1 v/v (20 mL, 1 equiv.) was added slowly and I the mixture was stirred for 30 min. Methyl bromoacetate (1.9 mL) was subsequently added and the mixture was stirred for 30 minutes at room temperature. The mixture was diluted with cethyl acetate and quenched with a saturated aquous ammonium chloride solution. The organic llayer was washed with water and brine, dried over sodium sulfate, filtered and evaporated in vacuo. The residue was purified by chromatography on sisilica gel (eluent:

TLC: Rf= 0.38, ethyl acetate/heptane 3/1 v/v on silica.

15 (c) BzlSO₂-norVal(cyclo)-Gly-OMe.

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Cbz-norVal(cyclo)-Gly-OMe (4.7 g) was dissolved in 40 mL of methanool, 500 mg 10% palladium on charcoal was added, 7.4 mL of a 2N hydrochloric acid was added and hydrogenated at atmospheric pressure for 1 hour at room temperature. The reaction mixture was filtered, evaporated in vacuo and immediately used in the next step as H-noorVal(cyclo)-Gly-OMe. HCl.

The crude amine was dissolved in dichloromethane (50 mL) and benzylsulfonyylchloride (2.82 g) was added slowly at 0 °C. Triethylamine was added to keep the pH at 8 durings the reaction. The mixture was stirred for 1 hour at room temperature, whereafter the mixture was washed with water and brine, dried over sodium sulfate, filtered and evaporated in vacuo. The residue was purified by chromatography on silica gel (eluent: dichloromethane/methanol 95/5 v/v) to yield BzlSO₂-norVal(cyclo)-Gly-OMe (2 g)

TLC: Rf=0.87, ethyl acetate/pyridine/acetic acid/water=63/20/6/11 v/v/v/v on silica.

(d) BzlSO₂-norVal(cyclo)-Gly-OH.

The saponification of BzlSO₂-norVal(cyclo)-Gly-OMe (2 g) was done according to the procedure described in Example 1. Yield: 1.8 g.

TLC: Rf=0.40, ethyl acetate/pyridine/acetic acid/water=63/20/6/11 v/v/v/v on i silica.

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(e) BzlSO₂-norVal(cyclo)-Gly-LysΨ[COCO]-OH

Coupling between BzlSO₂-norVal(cyclo)-Gly-OH and H-Lys(Boc)Y[CHOHCO]-OMe.HCl, saponification, oxidation, deprotection and purification were done according to procedures described in Example 1. Yield: 107 mg of the title compound.

5 Rt (LC): 24.45 min. 20% A/80% B to 20% A/20% B/60% C in 40 min.

Example 9.

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EthylSO₂-D-Cha-Pro-LysΨ[COCO]-O-iPropyl

H-Lys(Boc)Ψ[CHOHCO]-O-i-Propyl.HCl (0.32 g) was prepared using the procedure described for H-Lys(Boc)Ψ[CHOHCO]-OEt.HCl in Example 3 starting from

Cbz-Lys(Boc)Ψ[CHOHCO]-OMe (0.49 g) and 2-propanol. The DCC/HOBt-ccoupling between EthylSO₂-D-Cha-Pro-OH (239mg) and H-Lys(Boc)Ψ[CHOHCO]-O-i-Propyyl.HCl (316 mg), Dess-Martin oxidation, deprotection using trifluoroacetic acid and purification were done according to the procedures described in Example 1. Yield: 123 mg of the titlee compound.

Rt (LC): 43.0 min. 20% A/80% B to 20% A/20% B/60% C in 40 min. and maintain this mixture of eluens for an additional 10 min.

Example 10.

BzlSO2-norVal(cyclo)-Gly-LysY[COCO]-Azetidine

The procedures described in Example 1 were used to prepare the title compound. Cbz-Lys(Boc)Ψ[CHOHCO]-Azetidine (2.26 g) was prepared from Cbz-Lys(Boc)Ψ[CHOHCO]-OH (2.7 g) as described for Cbz-Lys(Boc)Ψ[CHOHCO]-NHBzl. Hydrogeenation of Cbz-Lys(Boc)Ψ[CHOHCO]-Azetidine (269 mg) yielded H-Lys(Boc)Ψ[CHOHCO]-Azetidine.HCl (214 mg). Then DCC/HOBt-coupling between BzlSO₂-norVal(cyclo)-Gly-OHI (175 mg) and H-Lys(Boc)Ψ[CHOHCO]-Azetidine.HCl (214 mg), Dess-Martin oxidation, ddeprotection using trifluoroacetic acid and purification yielded 84 mg of the title compound.

Rt (LC): 27.8 min. 20% A/80% B to 20% A/20% B/60% C in 40 min.

Example 11.

30 BzlSO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-Azetidine

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Cbz-Lys(Boc)Y[CHOHCO]-Azetidine was prepared according to procedures described in Example 10. The hydrogenation, coupling to BzlSO₂-norLeu(cyclo)-Glyy-OH, oxidation, deprotection and purification were also done according to procedures described in Example 1. Yield: 100 mg of the title compound.

5 Rt (LC): 33.61 min. 20% A/80% B to 20% A/20% B/60% C in 40 min.

Example 12.

BzISO₂-norLeu(cyclo)-Gly-Lys(Ethoxycarbonyl)Ψ[COCO]-Azetidine

- (a) BzISO2-norLeu(cyclo)-Gly-LysY[CHOHCO]-Azetidine.TFA
- BzlSO₂-norLeu(cyclo)-Gly-Lys(Boc)Ψ[CHOHCO]-Azetidine (prepared according to procedures described in Example 10) (220 mg) was dissolved in 10 mL of dichloromethane/trifluoroacetic acid 1/1 v/v and stirred for 2 hours at room temperature. Solvents were removed by evaporation and the residue titruated with diethyl ether. Yield: 267 mg.
- 15 TLC: Rf=0.57, ethyl acetate/pyridine/acetic acid/water=63/20/6/11 v/v/v/v on s silica.
 - (b) BzlSO2-norLeu(cyclo)-Gly-Lys(Ethoxycarbonyl)Y[CHOHCO]-Azetidine

BzlSO₂-norLeu(cyclo)-Gly-LysΨ[CHOHCO]-Azetidine.TFA (267 mg) was dilissolved in 10 mL of N,N-dimethylformamide and 46 μL of ethylchloroformate was added after which the pH was adjusted to 8.5 with triethylamine. After stirring for 16 hours at room temperature, the reaction mixure was diluted with ethyl acetate, washed with water, 5% sodium hydrogencarbonate, 2% citric acid and brine, dried on sodium sulfate, filtered and evaporated to drynesss to yield 150 mg of the title compound.

TLC: Rf= 0.53, dichloromethane/methanol 9/1 v/v on silica.

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(c) BzISO₂-norLeu(cyclo)-Gly-Lys(Ethoxycarbonyl)Ψ[COCO]-Azetidine

Oxidation and purification of BzlSO₂-norLeu(cyclo)-Gly-Lys(Ethoxycarbonnyl)Ψ[CHOHCO]-Azetidine (150 mg) were done according to procedures described in Example : 1. Yield 25 mg. Rt (LC): 26.42 min. 20% A/60% B/20%C to 100% C in 40 min.

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Example 13.

BzlSO2-norLeu(cyclo)-Gly-LysΨ[COCO]-O-iPropyl

Coupling between BzlSO₂-norLeu(cyclo)-Gly-OH (described in Example 1) and H-Lys(Boc)\(P(CHOHCO)\)-O-iPropyl.HCl (described in Example 9), oxidation, deprotection and purification were done according to procedures described in Example 1. Yieldd: 400 mg of the title compound.

5 Rt (LC): 40 min. 20% A/80% B to 20% A/20% B/60% C in 40 min.

Example 14.

BzlSO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-NH-iPropyl

- (a) BzlSO₂-norLeu(cyclo)-Gly-Lys(Boc)Ψ[CHOHCO]-OH
- The DCC/HOBt-coupling between 1.96 g of BzlSO₂-norLeu(cyclo)-Gly-OH ε and 2.20 g of H-Lys(Boc)Ψ[CHOHCO]-OMe.HCl and saponification of the product were perfformed according to the procedures described in example 1. Yield: 3.1 g of the crude title compound.

 TLC: Rf=0.4, ethyl acetate/pyridine/acetic acid/water=66/20/6/11 v/v/v/v on sililica.
- 15 (b) BzISO₂-norLeu(cyclo)-Gly-Lys(Boc)Ψ[CHOHCO]-NH-iPropyl

The EDCI/HOBt-coupling between 0.4 mmol BzlSO₂-ncorLeu(cyclo)-Gly-Lys(Boc)\P[CHOHCO]-OH and 0.105 mL of isopropylamine, Dess Martin oxidation (reaction time: 19 h) and deprotection were done according to the procedures describbed in example 1. Yield: 150 mg of the title compound.

20 Rt(LC): 34.01 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

Example 15.

BzlSO2-norLeu(cyclo)-Gly-LysY[COCO]-NH-nPropyl

The EDCI/HOBt-coupling between 0.4 mmol BzlSO₂-noorLeu(cyclo)-Gly-Lys(Boc)Ψ[CHOHCO]-OH and 0.101 mL of propylamine, Dess Martin oxidation (reaction time: 24 h) and deprotection were done according to the procedures described in example 1. Yield: 144 mg of the title compound.

Rt(LC): 34.22 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

30 **Example 16.**

BzISO2-norLeu(cyclo)-Gly-LysΨ[COCO]-NH-Methyl

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The EDCI/HOBt-coupling between 0.4 mmol BzlSO₂-noorLeu(cyclo)-Gly-Lys(Boc)Y[CHOHCO]-OH and methylamine (0.4 mL of a 3 M scolution in N,N-dimethylformamide), Dess Martin oxidation (reaction time: 20 h) and deprotection were done according to the procedures described in example 1. Yield: 127 mg of the title compound.

5 Rt(LC): 28.36 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

Example 17.

BzISO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-pyrrolidinyl

The EDCI/HOBt-coupling between 0.4 mmol BzISO₂-neorLeu(cyclo)-Gly-10 Lys(Boc)Ψ[CHOHCO]-OH and 0.102 mL of pyrrolidine, Dess Martin oxidation (reaction time: 14 days) and deprotection were done according to the procedures described in example 1. Yield: 125 mg of the title compound.

Rt(LC): 36.87 and 37.38 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 400 min.

15 **Example 18.**

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BzlSO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-N-Ethyl

The EDCI/HOBt-coupling between 0.4 mmol BzISO₂-neorLeu(cyclo)-Gly-Lys(Boc)\(Partial (CHOHCO)\)-OH and ethylamine (1.78 mL of a 0.7 M ssolution in N,N-dimethylformamide), Dess Martin oxidation (reaction time: 20 h) and deprotection were done according to the procedures described in example 1. Yield: 115 mg of the title: compound.

Rt(LC): 31.30 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

Example 19.

BzlSO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-morpholin-4-yl

25 The EDCI/HOBt-coupling between 0.4 mmol BzlSO₂-nnorLeu(cyclo)-Gly-Lys(Boc)Ψ[CHOHCO]-OH and 0.107 mL of morpholine, Dess Martin ooxidation (reaction time: 6.5 days) and deprotection were done according to the procedures described in example 1. Yield: 148 mg of the title compound.

Rt(LC): 33.73 and 34.17 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 400 min.

Example 20.

BzlSO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-(1,1-dioxo)thiomorpholin-4-yl

To a solution of 2.47 g of thiomorpholine in 25 mL of methanol was added \$5.75 g of di-tert-butyl dicarbonate and 4 mL of triethylamine. After stirring at room temperatuure for 3h, 50 mL of ethyl acetate was added and this solution was washed with water adjusted to pH 3 with hydrochloric acid, water, aqueous 5% sodium hydrogencarbonate and bbrine, dried over magnesium sulfate and concentrated to give 4.73 g of N-tert-butyloxycarbonyl thiomorpholine. This residue (4.73 g) was dissolved in 50 mL of dichloromethane and 50 nmL of water was added. To this stirred mixture was added 11 g 3-chloroperoxybenzoic acid (800 - 90% purity) in small portions keeping the reaction mixture at pH 7. After stirring at room temperature for 16 h the water layer was separated, the organic layer washed with 5% aqueous soodium thiosulfate, 5% aqueous sodium hydrogencarbonate (three times) and brine, dried over magnesium sulfate and concentrated. The residue was purified by chromatography on silica [gel (eluent: ethyl acetate / heptanes 2/3 v/v) to give 5.7 g of N-tert-butyloxycarbonyl thiomorpholine 1,1-dioxide. This sulfon (0.625 g) was dissolved in 50 mL of a 3M hydrogenchloride solution in dioxane and after stirring for 4 hours at room temperature the reaction mixture was conncentrated to give 0.579 g of thiomorpholine 1,1-dioxide hydrochloride.

The EDCI/HOBt-coupling between 0.4 mmol BzISO₂-ncorLeu(cyclo)-Gly-Lys(Boc)Ψ[CHOHCO]-OH and 0.21 g of thiomorpholine 1,1-dioxide hyddrochloride, Dess Martin oxidation (reaction time: 3 days) and deprotection were done εaccording to the procedures described in example 1. Yield: 180 mg of the title compound.

20 Rt(LC): 33.64 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

Example 21.

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BzlSO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-N(Methyl)(Methoxy)

The EDCI/HOBt-coupling between 0.4 mmol BzISO₂-nαorLeu(cyclo)-Gly-Lys(Boc)Ψ[CHOHCO]-OH and 0.12 g of N,O-dimethylhydroxylamine, Desss Martin oxidation (reaction time: 3.5 days) and deprotection were done according to the proceedures described in example 1. Yield: 136 mg of the title compound.

Rt(LC): 33.80 and 34.53 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 400 min.

30 Example 22.

BzlSO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-(2-(carboxamid)azetidin-1-yl)

The DCC/HOBt-coupling between 1.13 g N-tert-butyloxycarbonyl-L-azettidine-2-carboxylic acid and 1.38 g ammmonium chloride was performed as described in examplee 1 to give 0.468 g

23

of N-tert-butyloxycarbonyl-L-azetidine-2-carboxamide. This amide (0.224 g) was dissolved in 5 mL of a 3M hydrogenchloride solution in dioxane. After stirring for 33 hours at room temperature the reaction mixture was concentrated to give 0.17 g of azetidine-2-carboxamide hydrochloride.

5 The EDCI/HOBt-coupling between 0.4 mmol BzlSO₂-noorLeu(cyclo)-Gly-Lys(Boc)Ψ[CHOHCO]-OH and 0.17 g of azetidine-2-carboxamide hydrochlopride, Dess Martin oxidation (reaction time: 20 h) and deprotection were done according to the procedures described in example 1. Yield: 58 mg of the title compound.

Rt(LC): 26.43 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

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Example 23.

nPropylSO₂-D-Cha-Pro-LysΨ[COCO]-O-iPropyl

(a) Boc-D-Cha-Pro-OBzl

To a stirred solution of 11.64 g of Boc-D-Cha-OH in 100 mL of dichloromeethane at 0°C was added 6.36 g of HOBt and 9.72 g of DCC. After 20 minutes a solution of 110.35 g of H-Pro-OBzl. HCl in 40 mL of dichloromethane adjusted with N,N-diisopropyl ethylaamine to pH 8 was added. After 16 h the reaction mixture was filtered and the filtrate was washed I successively with water, 0.1 N hydrochloric acid, water, aqueous 5% sodium hydrogencarbonaate and brine. All aqueous washes were extracted twice with ethyl acetate, all organic extractss combined, dried over sodium sulfate and concentrated. To the residue was added a mixturee of ethyl acetate/heptanes = 1/1 (v/v), the resulting suspension filtered and the filtrate purified by chromatography on silica gel (eluent: ethyl acetate/heptanes = 1/1 v/v) to yieldd 19.34 g of Boc-D-Cha-Pro-OBzl.

TLC: Rf=0.8, dichloromethane/methanol=9/1 v/v on silica.

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(b) nPropylSO2-D-Cha-Pro-OBzl

Boc-D-Cha-Pro-OBzl (1.01 g) was dissolved in 42 mL of a 3M hydrogenchhloride solution in dioxane. After stirring for 2 hours at room temperature the reaction mixture was concentrated. The residue was dissolved in 35 mL of dichloromethane and cooled to 0°°C. To this stirred solution was added 0.22 mL of 1-propanesulfonyl chloride and the pH adjuusted to 8.5. After stirring for 24 h at room temperature the reaction mixture was concentrated. The residue was dissolved in ethyl acetate, washed successively with aqueous 5% sodium hydrogencarbonate, water, aqueous 5% citric acid and brine, dried over magnesium sulfate and concentrated. The

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crude product was purified by chromatography on silica gel (eluent: ethyl accetate/ heptanes = 1/1 v/v) to yield 0.85 g of nPropylSO₂-D-Cha-Pro-OBzl.

TLC: Rf=0.6, ethyl acetate/ heptanes = 1/1 v/v on silica.

5 (c) nPropylSO₂-D-Cha-Pro-LysΨ[COCO]-O-iPropyl

nPropylSO₂-D-Cha-Pro-OBzl (0.85 g) was hydrogenated using the proceddure described in example 1 to give 0.54 g of nPropylSO₂-D-Cha-Pro-OH. The DCC/HOBt cooupling of 225 mg of nPropylSO₂-D-Cha-Pro-OH and of H-Lys(Boc)Ψ[CHOHCO]-O-iPropyyl.HCl and Dess Martin oxidation were performed according to the procedures described in ι example 9. The Boc-group was removed using a 3M hydrogenchloride solution in dioxane ass described above and the crude product purified using the preparative HPLC method describbed in example 1. Yield: 47 mg of the title compound.

Rt(LC): 27.6 min. 20% A/ 60% B/ 20% C to 20% A/ 80% C in 30 min, then to 100% C in 10 min.

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Example 24.

(10-Camphor)SO2-D-Cha-Pro-LysY[COCO]-O-iPropyl

The title compound was prepared from Boc-D-Cha-Pro-OBzl and (-)-1(0-camphorsulfonyl chloride using the procedures described in example 23. Yield: 12% from Boc-D-Cha-Pro-OBzl. Rt(LC): 33.6 min. 20% A/ 60% B/ 20% C to 20% A/ 80% C in 30 min, thenn to 100% C in 10 min.

Example 25.

PhenylSO2-D-Cha-Pro-LysY[COCO]-O-iPropyl

The title compound was prepared from Boc-D-Cha-Pro-OBzl and benzennesulfonyl chloride using the procedures described in example 23. Yield: 9% from Boc-D-Cha-Pro-OBzl. Rt(LC): 29.3 min. 20% A/ 60% B/ 20% C to 20% A/ 80% C in 30 min, thern to 100% C in 10 min.

30 Example 26.

MethylSO₂-D-Cha-Pro-LysΨ[COCO]-O-iPropyl

The title compound was prepared from Boc-D-Cha-Pro-OBzl and methannesulfonyl chloride using the procedures described in example 23. Yield: 18% from Boc-D-Cha-l-Pro-OBzl.

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Rt(LC): 24.3 min. 20% A/ 60% B/ 20% C to 20% A/ 80% C in 30 min, then 1 to 100% C in 10 min.

Example 27.

5 <u>iPropylSO₂-D-Cha-Pro-LysΨ[COCO]-O-iPropyl</u>

The title compound was prepared from Boc-D-Cha-Pro-OBzl and isopropyl/Isulfonyl chloride using the procedures described in example 23. Yield: 2% from Boc-D-Cha-Proo-OBzl. Rt(LC): 26.8 min. 20% A/ 60% B/ 20% C to 20% A/ 80% C in 30 min, then to 100% C in 10 min.

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Example 28.

BenzylSO₂-D-Cha-Pro-LysΨ[COCO]-O-iPropyl

The title compound was prepared from Boc-D-Cha-Pro-OBzl and α-tolueneesulfonyl chloride using the procedures described in example 23. Yield: 11% from Boc-D-Cha-P?ro-OBzl.

15 Rt(LC): 30.4 min. 20% A/ 60% B/ 20% C to 20% A/ 80% C in 30 min, then 1 to 100% C in 10 min.

Example 29.

nButylSO₂-D-Cha-Pro-LysΨ[COCO]-O-iPropyl

The title compound was prepared from Boc-D-Cha-Pro-OBzl and 1-butangesulfonyl chloride using the procedures described in example 23. Yield: 29% from Boc-D-Cha-PPro-OBzl.

Rt(LC): 29.3 min. 20% A/ 60% B/ 20% C to 20% A/ 80% C in 30 min, thenn to 100% C in 10 min.

25 Example 30.

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[3-(benzylsulfonylamino)-6-methyl-2-oxo-1,2-dihydropyridinyl]-acetyl-Lys \(\mathbb{I}\)[(COCO]-O-iPropyl The DCC/HOBt coupling of 151 mg of [3-(benzylsulfonylamino)-6--methyl-2-oxo-1,2-dihydropyridinyl]-acetic acid (WO 97/01338) and 205 mg of H-Lys(Boc)\(\mathbb{Y}\)[CHOHCO]-O-iPropyl.HCl, Dess Martin oxidation, deprotection and purification were performed according to the procedures described in example 9 to give 91 mg of the title compound.

Rt(LC): 34.7 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

Example 31.

[3-(benzylsulfonylamino)-2-oxo-1,2-dihydropyridinyl]-acetyl-LysΨ[COCO]-O-iPropyl
The DCC/HOBt coupling of 178 mg of [3-(benzylsulfonylamino)-2-oxo-1,2-ddihydropyridinyl]acetic acid (WO 97/46207) and H-Lys(Boc)Ψ[CHOHCO]-O-iPropyl.HCl annd Dess Martin
oxidation were performed according to the procedures described in examplee 9. Deprotection
using hydrogenchloride in dioxane and purification were performed according to the procedures
described in example 23 to give 116 mg of the title compound.

Rt(LC): 32.4 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

10 Example 32.

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[3-(benzylsulfonylamino)-6-methyl-2-oxo-1,2-dihydropyridinyl]-acetyl-LysΨ[CCOCO]-NH₂
The DCC/HOBt coupling of 286 mg of [3-(benzylsulfonylamino)-6-nmethyl-2-oxo-1,2-dihydropyridinyl]-acetic acid (WO 97/01338) and H-Lys(Boc)Ψ[CHOHCO]-OMe HCl, according to the procedure described in example 1 yielded 0.51 g of [3-(benzzylsulfonylamino)-6-methyl-2-oxo-1,2-dihydropyridinyl]-acetyl-LysΨ[CHOHCO]-OMe. Saponiification of this methyl ester, EDCI/HOBt coupling with ammonium chloride, Dess Martin oxidation, deprotection and purification were performed according to the procedures desscribed in example 14 to give 76 mg of the title compound.

Rt(LC): 26.9 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

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Example 33.

BzlSO₂-Aad-Pro-LysΨ[COCO]-OH

(a) BzlSO2-Aad(OtBu)-OH

To a stirred solution of 0.5 g of H-Aad(OtBu)-OH in 4.4 mL of aqueous 1 N is sodium hydroxide was added 0.42 g of benzylsulfonylchloride in 2 mL of dioxane. After 116 hours at room temperature, additional 1.4 mL of aqueous 2 N sodium hydroxide, 0.5 mL off dioxane and 0.09 g of benzylsulfonylchloride were added and the reaction mixture stirred for an additional day. The dioxane was removed, water was added, the mixture made acid (pH 3) tusing hydrochloric acid and extracted twice with diethyl ether. The combined ether layers were dried over sodium sulfate and concentrated to give 235 mg of BzlSO₂-Aad(OtBu)-OH.

TLC: Rf=0.7, dichloromethane/methanol/water = 14/6/1 v/v/v on silica.

(b) BzlSO2-Aad(OtBu)-Pro-OH

DCC/HOBt coupling of 235 mg of BzlSO₂-Aad(OtBu)-OH and 168 mg of 1H-Pro-OBzl.HCl followed by hydrogenation as described in example 23 yielded 193 mg of the tittle compound. TLC: Rf=0.6, ethyl acetate/pyridine/acetic acid/water=163/20/6/11 v/v/v/v on sisilica.

(c) BzlSO₂-Aad-Pro-LysΨ[COCO]-OH

The DCC/HOBt coupling of 193 mg of BzlSO₂-Aad(OtBu)-Proo-OH and H-Lys(Boc)Ψ[CHOHCO]-OMe HCl, saponification, Dess Martin oxidation, edeprotection and purification were performed according to the procedures described in example 132 to give 85 mg of the title compound.

Rt(LC): 26.1 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

Example 34.

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BzlSO₂-Glu-Pro-LysΨ[COCO]-OH

Starting with H-Glu(OtBu)-OH according to the route described in example 33 gave the title compound. Yield: 3% from H-Glu(OtBu)-OH.

Rt(LC): 22.6 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

Example 35.

20 BzlSO₂-Asp-Pro-LysΨ[COCO]-OH

Starting with H-Asp(OtBu)-OH according to the route described in example: 33 gave the title compound. Yield: 18% from H-Asp(OtBu)-OH.

Rt(LC): 21.9 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

25 Example 36.

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EtSO₂-D-Tyr(Me)-Pro -LysΨ[COCO]-NH₂

(a) EtSO₂-D-Tyr(Me)-Pro-OH

DCC/HOBt coupling of 2.22 g of Boc-D-Tyr(Me)-OH and 2.0 g of H-Pro-OB3zl . HCl, removal of the Boc protecting group, sulfonylation using ethane sulfonyl chloride and I hydrogenation of the benzyl ester using the procedures described in example 23 yielded 11.0 g of the title compound

TLC: Rf=0.23, dichloromethane/methanol = 95/5 v/v on silica.

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(b) EtSO2-D-Tyr(Me)-Pro -LysY[COCO]-NH2

The DCC/HOBt coupling of 254 mg of EtSO₂-D-Tyr(Me)-Proo-OH and H-Lys(Boc)\P[CHOHCO]-OMe.HCl, saponification, EDCI/HOBt coupling with ammonium chloride, Dess Martin oxidation, deprotection and purification were performed 1 according to the procedures described in example 32 to give 83 mg of the title compound.

Rt(LC): 28.0 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

Example 37.

EtSO₂-D-Tyr(Me)-Pro -LysΨ[COCO]-O-iPropyl

The DCC/HOBt coupling of 0.51 g of EtSO₂-D-Tyr(Me)-Proo-OH and H-Lys(Boc)Ψ[CHOHCO]-O-iPropyl.HCl, Dess Martin oxidation, deprotection and purification were performed according to the procedures described in example 9 to give 2223 mg of the title compound.

Rt(LC): 36.5 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

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Example 38.

EtSO₂-D-Tyr(Me)-Pro -LysΨ[COCO]-Azetidine

The DCC/HOBt coupling of 307 mg of EtSO₂-D-Tyr(Me)-Proo-OH and H-Lys(Boc)Ψ[CHOHCO]-Azetidine.HCl, Dess Martin oxidation, deprotection and purification were performed according to the procedures described in example 10 to give 183 mg of the title compound.

Rt(LC): 36.4 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

Example 39.

25 <u>EtSO₂-D-Tyr(Me)-Pro -LysΨ[COCO]-N-(4-chloropropyl)</u>

The title compound (43 mg) was obtained as second product in the purification of example 38. Rt(LC): 38.1 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 min.

Example 40.

30 BzlSO₂-D-Dpa-Pro-LysΨ[COCO]-O-iPropyl

(a) BzlSO₂-D-Dpa-Pro-OH

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Removal of the Boc group of 1.5 g of Boc-D-Dpa-Pro-OBzl (WO 97/319337), reaction with benzylsulfonyl chloride and removal of the benzyl ester according to the proceddures described in example 23 to yield 1.0 g of the title compound.

TLC: Rf=0.63, ethyl acetate/pyridine/acetic acid/water=163/20/6/11 v/v/v/v on 1 silica.

(b) BzlSO₂-D-Dpa-Pro-LysΨ[COCO]-O-iPropyl

The DCC/HOBt coupling of 0.31 g of BzlSO₂-D-Dpa-Pro-OH and H-Lys(Booc)Ψ[CHOHCO]-O-iPropyl.HCl, Dess Martin oxidation, deprotection and purification were perfformed according to the procedures described in example 9 to give 50 mg of the title compound.

10 Rt(LC): 32.2 min. 20% A/ 60% B/ 20%C to 20% A/ 80% C in 30 min, then to 100% C in 10 min.

Example 41.

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EtSO₂-Leu-Pro-LysΨ[COCO]-O-iPropyl

15 (a). EtSO₂-Leu-OMe

A stirred solution of 3.0 g of H-Leu-OMe.HCl in 30 mL of dichloromethane wwas adjusted to pH 8 using triethylamine and cooled at 0°C. Then 3.2 mL of ethanesulfonyl chloride and 2.3 mL of triethylamine were added. After stirring for 16 h at room temperature the reaction mixture was washed successively with 0.5 N hydrochloric acid, water and aquecous 5% sodium hydrogencarbonate and concentrated. The crude product was purified by chhromatography on silica gel (eluent: dichloromethane/ methanol = 9/1 v/v) to yield 3.3 g of EtSO₇₂-Leu-OMe.

TLC: Rf=0.69, dichloromethane/ ethyl acetate = 9/1 v/v on silica.

(b). EtSO2-Leu-Pro-OH

EtSO₂-Leu-OMe (3.3 g) was saponified (procedure example 1), coupled with H-Pro-OBzl (procedure example 23) and the resulting dipetide was hydrogenated (proceedure example 23) using the indicated procedures to give 3.4 g of the title compound.

TLC: Rf=0.11, dichloromethane/ ethyl acetate = 9/1 v/v on silica.

30 (c). EtSO₂-Leu-Pro-LysΨ[COCO]-O-iPropyl

The DCC/HOBt coupling of 145 mg of EtSO₂-Leu-Pro-OH and H-Lys(Boc))Ψ[CHOHCO]-O-iPropyl.HCl, Dess Martin oxidation, deprotection and purification were performed according to the procedures described in example 23 to give 120 mg of the title compound.l.

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Rt(LC): 16.6 min. 20% A/60% B/20%C to 20% A/80% C in 30 min.

Example 42.

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BzlSO₂-norLeu(cyclo)-Gly-Acg Ψ[COCO]-Azetidine

(a). H-Acg(Boc)[CHOHCO]-OMe.HCl

To a solution of 3-[4-(1,1-dimethylethoxycarbonylamino)cyclohexyl]-2-hydrcoxy-3-nitro-propionic acid methyl ester (Lyle et al, Bioorg. Med. Chem. Lett., 7, 67-72 ([1997]) (294 mg) in methanol (100 mL) was added 2N hydrochloric acid (0.425 mL) and 10 % paalladium on activated carbon powder (0.45 g) and this suspension was hydrogenated at attmospheric pressure at room temperature for 16 hours. The palladium catalyst was removed by filtration and the solvent was removed by evaporation at reduced pressure yielding H-· Acg(Boc)Ψ[CHOHCO]-OMe.HCl (289 mg) as a mixture of diastereomers.

TLC: R_f= 0.26, silica gel, ethyl acetate/pyridine/acetic acid/water=232/31/18//7 v/v/v/v.

15 (b) BzlSO₂-norLeu(cyclo)-Gly-Acg Ψ[COCO]-Azetidine

The DCC/HOBt-coupling between 0.27 g of BzlSO₂-norLeu(cyclo)-Gly-OH i and 0.25 g of H-Acg(Boc)Ψ[CHOHCO]-OMe.HCl, saponification, EDCI/HOBt-coupling with azetidine hydrochloride, Dess Martin oxidation and deprotection were done according to the procedures described in example 1. Yield: 82 mg of the title compound.

20 Rt(LC): 34.8 and 35.4 min. 20% A/ 80% B to 20% A/ 20% B/ 60% C in 40 rmin.

Example 43.

EthylSO₂-D-Cha-Pro-AcgΨ[COCO]-OiPropyl

(a) Cbz-Acg(Boc)Ψ[CHOHCO]-OMe

A stirred solution of 0.34 g of H-Acg(Boc)Ψ[CHOHCO]-OMe.HCl in 10 mL of acetonitrile and 10 mL of N,N-dimethylformamide is adjusted to pH 8 using N,N-diisoprropylethylamine. To this solution 0.24 g of N-benzyloxycarbonyloxysuccinimide was added. After stirring at room temperature for one hour the reaction mixture was concentrated. The residue dissolved in ethyl acetate, washed with water and brine, dried over sodium sulfate and concentrated. The residue was purified by chromatography on silica gel (eluent: ethyl acetate / heptannes 2/3 v/v) to give 0.287 mg of Cbz-Acg(Boc)Ψ[CHOHCO]-OMe.

TLC: Rf=0.25, ethyl acetate / heptanes 1/1 v/v on silica.

(b) Cbz-Acg(Boc)Ψ[CHOHCO]-OiPropyl

To a stirred mixture of 5 mL of tetrahydrofuran and 1 mL of 2-propanol tunder a nitrogen atmosphere was added slowly added 2.5 mL of a 1.6N n-butyllithium solution i in hexanes. After 20 minutes a solution of 0.28 g of Cbz-Acg(Boc)Ψ[CHOHCO]-OMe in 5mL obf 2-propanol was added and stirred for 2 h at room temperature. Then 0.5 mL of acetic acid was added and the reaction mixture was concentrated. The residue dissolved in ethyl acetate, waashed with water, dried over sodium sulfate and concentrated. The residue was purified by chnromatography on silica gel (eluent: ethyl acetate / heptanes 2/3 v/v) to give 0.2233 mg of Cbz-Acg(Boc)Ψ[CHOHCO]-OiPropyl.

TLC: Rf=0.25, ethyl acetate / heptanes 1/2 v/v on silica.

(b) EthylSO₂-D-Cha-Pro-AcgΨ[COCO]-OiPropyl

To a solution of 0.22 g of Cbz-Acg(Boc)Ψ[CHOHCO]-OiPropyl in N,N-dimetthylformamide were added 10% palladium on activated carbon (80 mg) and 2M hydrochloric aacid (0.23 mL) and this suspension was hydrogenated at atmospheric pressure for 1 hour at rooom temperature. The palladium catalyst was removed by filtration. This fitrate was used in a DCCC/HOBt coupling with 0.166 g of EthylSO₂-D-Cha-Pro-OH using the procedure describbed in example 1. The product was oxidised using the Dess Martin reagent, the Boc-group removed and purified using the procedures described in example 1. Yield: 100 mg of the title compound. Rt(LC): 30.0 min. 20% A/ 60% B/ 20% C to 20% A/ 80% C in 30 min

Example 44.

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Preparation of EtSO₂-B-X-LysΨ[COCO]-O-iPropyl derivatives on solid phasee.

25 (a) Teoc-Lys(Boc)Ψ[CHOHCO]-O-iPropyl

Cbz-Lys(Boc)Ψ[CHOHCO]-OMe (10 g) was hydrogenated under the conditions described in example 1f to afford H-Lys(Boc)Ψ[CHOHCO]-OMe in quantitative yield. The crude product was treated with 2-(trimethylsilyl)ethoxycarbonyl hydroxy-succinimide ((6.7 g) in N,N-dimethylformamide (100 mL) in the presence of N,N-diisopropylethylaminee (pH = 8) for 2 hours at room temperature. The reaction mixture was evaporated to drynesss and the residue was dissolved in ethyl acetate and washed with 2% aqueous citric acid, wvater, 5% aqueous sodium hydrogencarbonate and brine. Drying over sodium sulfate and evaporaation of the solvent

afforded, after chromatography on silica gel (eluent: ethyl acetate/heptane = 1/1 v/v), Teoc-Lys(Boc)Ψ[CHOHCO]-OMe (9.1 g). Subsequent transesterification was accomplished by adding dropwise Teoc-Lys(Boc)Ψ[CHOHCO]-OMe (2.8 g) to a stirred mixxture of isopropyl alcohol (5.4 mL), THF (27.1 mL) and 1.6 M n-butyl lithium in hexane (113.6 mL) at room temperature. After 1 hour the reaction mixture was cooled to 0°C and glaciaal acetic acid (2.5 mL) was added. The reaction mixture was concentrated to a small volume and I diluted with ethyl acetate, washed with water (2x) and dried over sodium sulfate. Filtration annd removal of the solvent in vacuo gave the crude product. Chromatography on silica ggel (eluent: ethyl acetate/heptane = 1/1 v/v) afforded the title compound (2.9 g).

10 TLC: Rf=0.53, heptane/ethyl acetate 1/1 v/v on silica.

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(b) Teoc-Lys(CO-O-methyl-resin) [CHOHCO]-O-iPropyl

Teoc-Lys(Boc)Ψ[CHOHCO]-O-iPropyl (2.8 g) was dissolved diethyl ether ([36 mL) and paratoluene sulfonic acid (1.8 g) was added. After 2 hours at 30 °C the reaction mixture was evaporated and the residue was dried *in vacuo* to give Teoc-LysΨ[CHOHCO]-O-iPropyl.

To a suspension of 4.2 g of hydroxymethyl-resin (Bachem, 1.02 mmol//g) in 50 mL of acetonitrile/dichloromethane (1/1 v/v) and triethylamine (1.81 mL) was added N,N-disuccinimidyl carbonate (3.36 g). The suspension was shaken for 2 hhours at ambient temperature on an orbital shaker. The resin was filtered off and washed with 1 dichloromethane, acetonitrile and dichloromethane (three times each) and dried. Teoc-LyssΨ[CHOHCO]-O-iPropyl (see above) was dissolved in 50 mL of acetonitrile/dichloromethane (11/1 v/v). The pH of the solution was adjusted to 8 using triethylamine. This solution was added too the resin and the suspension was shaken for 16 hours at room temperature. The solvent was remmoved by filtration and the resin was washed according to the procedures described earlier. Afteer drying in vacuo, 5.43 g of Teoc-Lys(CO-O-methyl-resin)Ψ[CHOHCO]-O-iPropyl was obtaineed.

(c) H-Lys(CO-O-methyl-resin)Ψ[CHOHCO]-O-iPropyl

A suspension of 2.5 g of Teoc-Lys(CO-O-methyl-resin)Ψ[CHOHCO]-O-iPropyl in trifluoroacetic acid/dichloromethane (50 mL, 1/9 v/v) was shaken for 45 min at room temperature. The resin was thoroughly washed with dichloromethane and I dried under high vacuum to give H-Lys(CO-O-methyl-resin)Ψ[CHOHCO]-O-iPropyl (2.5 g)

(d) Boc-X-Lys(CO-O-methyl-resin) Y[CHOHCO]-O-iPropyl

H-Lys(CO-O-methyl-resin)Ψ[CHOHCO]-O-iPropyl was divided over 4 reactoors in portions of 500 mg. The resin was washed with a 1% solution of N,N-diisoproppylethylamine in dichloromethane/N,N-dimethylformamide (3/2 v/v) and dichloromethane (thnree times each). Next, 10 mL of dichloromethane/N,N-dimethylformamide (3/2 v/v) was addded to the resin followed by building block Boc-X-OH (139 mg Boc-D-leu-OH, 139 mg Boc-ILeu-OH, 148 mg Boc-Gln-OH or 159 mg Boc-Phe-OH), 2-(1H-Benzotriazol-1-yl)-1,1,3,3-tettramethyluronium tetrafluoroborate (TBTU, 193 mg) and N,N-diisopropylethylamine (105 μL)). The suspension was shaken for 90 min at room temperature, whereafter the solvent was remooved by filtration. The resin was washed with dichloromethane/N,N-dimethylformamide ((3/2 v/v), N,N-dimethylformamide and dichloromethane (three times each) and dried.

(e) H-X-Lys(CO-O-methyl-resin)Ψ[CHOHCO]-O-iPropyl

The Boc-group of the four different X-blocks was removed under the samme conditions as described for the deprotection of the Teoc-group (see example 44c) to give foour times 500 mg of H-X-Lys(CO-O-methyl-resin) Y[CHOHCO]-O-iPropyl. This resin (500 mgz) was distributed over 5 reaction vessels.

(f) EtSO₂-B-X-Lys(CO-O-methyl-resin) Y[CHOHCO]-O-iPropyl

The couplings of the second building block EtSO₂-B-OH (27.0 mg EtSO₂-AAsn-OH, 26.8 mg EtSO₂-D-Leu-OH, 30.8 mg EtSO₂-D-Phe-OH, 36.8 mg EtSO₂-Nal-OH and 322.4 mg EtSO₂-D-3-Tiq-OH, prepared according to the methods as described in example 41)) were performed under the same conditions as described in procedure (d), based on 100 mg resinn. After work-up, the 20 reaction vessels (resulting from 4 different X blocks and 5 different B bblocks) were dried in vacuo.

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(g) EtSO₂-B-X-Lys(CO-O-methyl-resin)Ψ[COCO]-O-iPropyl

EtSO₂-B-X-Lys(CO-O-methyl-resin)Ψ[CHOHCO]-O-iPropyl (100 mg) was swollen in a solution of 1-hydroxy-1,2-benziodoxol-3(1H)-one 1-oxide (0.18 M) in dimmethylsulfoxide (2 mL) and dichloromethane (0.2 mL). The reaction mixture was allowed to shhake overnight at room temperature, whereafter the solvent was removed by filtration. Subsequeent washing with dimethylsulfoxide and dichloromethane (three times each) afforded, after dryying, EtSO₂-B-X-Lys(CO-O-methyl-resin)Ψ[COCO]-O-iPropyl.

(h) EtSO₂-B-X-LysΨ[COCO]-O-iPropyl

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A solution of trifluoroacetic acid/thioanisole (2 mL, 10/1 v/v) was added1 to EtSO₂-B-X-Lys(CO-O-methyl-resin)Y[COCO]-O-iPropyl (100 mg) and the reaction mixxture was shaken for 4 hours at room temperature. The resin was filtered, washed with trifluoroacetic acid (three times) whereafter the filtrate was evaporated to dryness in vacuo. The residuee was rinsed with heptane (2 mL) and vigorously stirred whereafter the heptane layer wass decanted. This procedure was repeated twice. The crude product was dried and directity applied on a preparative Supelcosil C18DB column (21 x 250 mm) for purification, using the following conditions: Flow: 20 mL/min; Buffers A: aqueous trifluoroacetic acid 0.1 lM, B: water, C: acetonitrile/water 6/4 v/v; Gradient (depending on the polarity of the product) 1 3% A - 67% B - 30% C to 3% A - 52% B - 45% C in 40 min. UV-detection at 210 nm. The main peaks, corresponding to the desired compounds, were isolated and lyophilized to givee the purified end products as depicted in table 44.

Table 44: Characterization (retention time on reversed phase HPLC and M+HI peak in electrospray mass spectrometry) of EtSO₂-B-X-LysΨ[COCO]-O-iPropyl prepaared on Hydroxymethyl-resin. HPLC conditions: Flow: 1.0 mL/min; Buffers A: water, B: acetonitril/water (6/4 v/v), C: 0.5 M phosphate-buffer pH = 2.1; Gradient: 0 - → 45 min 65 % A/15 % B/20 % C → 0 % A/80 % B/20 % C. UV-detection at 210 nm.

}			В		
:	Asn	D-Leu	D-Phe	Nal	D-3-Tiq
EtSO2-B-D-Leu-	Rt = 17.07	Rt =	Rt = 30.89	Rt = 38.17	RRt = 33.54
D/L-	min	27.20 min	min	min	min
LysΨ[COCO]-O-	M+H =	M+H =	M+H =	M+H =	M+H =
iPropyl	536.4	535.6	569.4	619.6	581.4
EtSO ₂ -B-Leu-	Rt = 17.17	Rt =	Rt = 32.89	Rt = 37.79	RRt = 33.60
D/L	min	30.78 min	min	min	min
Lys Y[COCO]-O-	M+H =	M+H =	M+H =	M+H =	M+H =
iPropyl	536.4	535.6	569.4	619.6	581.4
EtSO ₂ -B-Gln-	Rt = 5.40	Rt =	Rt = 18.05	Rt = 28.44	RRt = 21.27
D/L-	min	15.74 min	min	min	min
Lys Y[COCO]-O-	M+H =	M+H =	M+H =	M+H =	M+H =
iPropyl	551.2	550.4	584.4	634.4	596.4
EtSO ₂ -B-Phe-	Rt = 20.75	Rt =	Rt = 35.18	Rt = 39.47	FRt = 35.92
D/Lc	min	33.33 min	min	min	min
LysY[COCO]-O-	M+H =	M+H =	M+H =	M+H =	M+H =
iPropyl	570.4	569.4	603.4	653.6	615.6

10 Example 45.

The following compounds can be prepared by using the methods of the present invention:

 $CF_{3}SO_{2}\text{-}D\text{-}Cha\text{-}Pro\text{-}Lys\Psi[COCO]\text{-}O\text{-}iPropyl}$

 $MeSO_2\text{-}D\text{-}Tyr(Me)\text{-}Pro\text{-}Lys\Psi[COCO]\text{-}O\text{-}iPropyl$

n-ButylSO₂-D-Tyr(Me)-Pro-Lys\(\Psi\)[COCO]-O-iPropyl

15 CF₃SO₂-D-Tyr(Me)-Pro-Lys\{COCO}-O-iPropyl

BzlSO₂-D-Tyr(Me)-Pro-LysΨ[COCO]-O-iPropyl

 $EtSO_2\text{-}D\text{-}(p\text{-}OEt\text{-}Phe)\text{-}Pro\text{-}Lys\Psi[COCO]\text{-}O\text{-}iPropyl}$

EtSO2-D-Nle-Pro-LysY[COCO]-O-iPropyl

 $EtSO_2\text{-}D\text{-}Cha\text{-}Azt\text{-}Lys\Psi[COCO]\text{-}O\text{-}iPropyl$

20 EtSO₂-D-Cha-(N-cyclopentyl-Gly)-LysΨ[COCO]-O-iPropyl

EtSO2-D-Cha-Val-LysY[COCO]-O-iPropyl

EtSO₂-D-Cha-Pec-LysΨ[COCO]-O-iPropyl

EtSO₂-D-Cha-(3,4-dehydro-Pro)-LysΨ[COCO]-O-iPropyl

EtSO₂-D-Cha-Pro-LysΨ[COCO]-Azetidine

MeSO₂-D-Cha-Pro-LysY[COCO]-Azetidine

5 n-ButylSO₂-D-Cha-Pro-LysΨ[COCO]-Azetidine

CF₃SO₂-D-Cha-Pro-LysΨ[COCO]-Azetidine

BzlSO₂-D-Cha-Pro-LysΨ[COCO]-Azetidine

[3-(BzlSO₂amino)-2-oxo-1,2-dihydropyridinyl]-acetyl-LysΨ[COCO]-Azetidinee

[3-(BzlSO₂amino)-6-methyl-2-oxo-1,2-dihydropyridinyl]-acetyl-LysΨ[COCO]]-Azetidine

10 MeSO₂-D-Cha-Pro-AcgY[COCO]-O-iPropyl

 $n\text{-}ButylSO_2\text{-}D\text{-}Cha\text{-}Pro\text{-}Acg\Psi[COCO]\text{-}O\text{-}iPropyl$

 $CF_3SO_2\text{-}D\text{-}Cha\text{-}Pro\text{-}Acg\Psi[COCO]\text{-}O\text{-}iPropyl$

BzlSO₂-D-Cha-Pro-AcgΨ[COCO]-O-iPropyl

 $EtSO_2\text{-}D\text{-}Tyr(Me)\text{-}Pro\text{-}Acg\Psi[COCO]\text{-}O\text{-}iPropyl$

15 MeSO₂-D-Tyr(Me)-Pro-AcgΨ[COCO]-O-iPropyl

 $n\text{-}ButylSO_2\text{-}D\text{-}Tyr(Me)\text{-}Pro\text{-}Acg\Psi[COCO]\text{-}O\text{-}iPropyl$

CF₃SO₂-D-Tyr(Me)-Pro-Acg\(\Psi\)[COCO]-O-iPropyl

BzlSO₂-D-Tyr(Me)-Pro-AcgΨ[COCO]-O-iPropyl

EtSO₂-D-Tyr(Et)-Pro-AcgΨ[COCO]-O-iPropyl

20 EtSO₂-D-Nle-Pro-AcgΨ[COCO]-O-iPropyl

EtSO₂-D-Cha-Azt-AcgΨ[COCO]-O-iPropyl

EtSO₂-D-Cha-(N-cyclopentyl-Gly)-AcgΨ[COCO]-O-iPropyl

EtSO₂-D-Cha-Val-AcgΨ[COCO]-O-iPropyl

EtSO₂-D-Cha-Pec-AcgΨ[COCO]-O-iPropyl

25 EtSO₂-D-Cha-(3,4-dehydro-Pro)-AcgΨ[COCO]-O-iPropyl

EtSO₂-D-Cha-Pro-AcgΨ[COCO]-Azetidine

EtSO₂-D-Tyr(Me)-Pro-AcgΨ[COCO]-Azetidine

 $EtSO_2\text{-}D\text{-}Tyr(Me)\text{-}Pro\text{-}Acg\Psi[COCO]\text{-}NH_2$

 $MeSO_{z}\text{-}D\text{-}Cha\text{-}Pro\text{-}Acg\Psi[COCO]\text{-}Azetidine$

n-ButylSO₂-D-Cha-Pro-AcgΨ[COCO]-Azetidine

 $CF_3SO_2\text{-}D\text{-}Cha\text{-}Pro\text{-}Acg\Psi[COCO]\text{-}Azetidine$

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BzlSO₂-D-Cha-Pro-AcgΨ[COCO]-Azetidine

 $3-(BzISO_2-amino)-1-carboxymethyl-pyridin-2-one-Acg\Psi[COCO]-O-iPropyl$

 $3\text{-}(BzlSO_2\text{-}amino)\text{-}1\text{-}carboxymethyl\text{-}pyridin\text{-}2\text{-}one\text{-}Acg}\Psi[COCO]\text{-}Azetidine$

3-(BzlSO₂-amino)-1-carboxymethyl-6-methyl-pyridin-2-one-AcgΨ[COCO]-O-iPropyl

3-(BzlSO₂-amino)-1-carboxymethyl-6-methyl-pyridin-2-one-AcgY[COCO]-Azzetidine

The biological activities of the compounds of the present invention were deterrmined by the following test methods.

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I. Anti-thrombin assay

Thrombin (Factor IIa) is a factor in the coagulation cascade.

The anti-thrombin activity of compounds of the present invention was assessedd by measuring spectrophotometrically the rate of hydrolysis of the chromogenic substrate s-22238 exterted by thrombin. This assay for anti-thrombin activity in a buffer system was used to ϵ assess the IC50value of a test compound.

Test medium:

Tromethamine-NaCl-polyethylene glycol 6000 (TNP) bouffer

Reference compound: I2581 (Kabi)

Vehicle: 20

TNP buffer.

Solubilisation can be assisted with dimethylsulfoxide, mnethanol, ethanol, acetonitrile or tert.-butyl alcohol which are without advverse effects in concentrations up to 2.5% in the final reaction mixture.:.

Technique

Reagents*

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1. Tromethamine-NaCl (TN) buffer

Composition of the buffer:

(50 mmol)) 6.057 g Tromethamine (Tris) 5.844 g (100 mmol)) NaCl

Water to

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The pH of the solution is adjusted to 7.4 at 37 °C with HCI (10 mmol·l⁻¹).

2. TNP buffer

Polyethylene glycol 6000 is dissolved in TN buffer too give a concentration of 3 $g \cdot l^{-1}$.

3. S-2238 solution

One vial S-2238 (25 mg; Kabi Diagnostica, Sweden)) is dissolved in 20 ml TN buffer to give a concentration of 1.25 mg·mml⁻¹ (2 mmol·l⁻¹).

4. Thrombin solution

Human thrombin (16 000 nKat·vial⁻¹; Centraal Laborratorium voor Bloedtransfusie, Amsterdam, The Netherlands) is disssolved in TNP buffer to give a stock solution of 835 nKat·ml⁻¹. Immediately before use this solution is diluted with TTNP buffer to give a concentration of 3.34 nKat·ml⁻¹.

All ingredients used are of analytical grade

For aqueous solutions ultrapure water (Milli-Q quuality) is used.

Preparation of test and reference compound solutions

The test and reference compounds are dissolved in Milli--Q water to give stock concentrations of 10^{-2} mol·l⁻¹. Each concentration i is stepwise diluted with the vehicle to give concentrations of 10^{-3} , 10^{-4} and 10^{-5} mol·l⁻¹. The dilutions, including the stock solution, are ussed in the assay (final concentrations in the reaction mixture: $3 \cdot 10^{-3}$; 10^{-3} ; 10^{-3} ; 10^{-4} ; 10^{-4} ; $3 \cdot 10^{-5}$; $3 \cdot 10^{-6}$ and 10^{-6} mol·l⁻¹, respectively).

Procedure

At room temperature 0.075 ml and 0.025 ml test compound or reference compound solutions or vehicle are alternately pipetted innto the wells of a microtiter plate and these solutions are diluted with 0.1115 ml and 0.0165 ml TNP buffer, respectively. An aliquot of 0.030 ml S-22238 solution is added to each well and the plate is pre-heated and pre-inncubated with shaking in an incubator (Amersham) for 10 min. at 37 °CC. Following pre-incubation the hydrolysis of S-2238 is started by addition of 0.030 ml thrombin solution to each well. The plate is incubated (vwith shaking for

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30 s) at 37 °C. Starting after 1 min of incubation, the absorbance of each sample at 405 nm is measured every 2 min. for a period oof 90 min. using a kinetic microtiter plate reader (Twinreader plus, Flow Laboratories).

All data are collected in an IBM personal computer usingg LOTUS-MEASURE. For each compound concentration (expressed in mol·l⁻¹ reaction mixture) and for the blank the absorbance is plotted versus the reaction time in min.

Evaluation of responses: For each final concentration the maximum absorbanace was calculated from the assay plot. The IC₅₀-value (final concentration, expressed in μmol·l⁻¹, ccausing 50 % inhibition of the maximum absorbance of the blank) was calculated using the loggit transformation analysis according to Hafner et al. (Arzneim.-Forsch./Drug Res. 1977, 27(II): 1871-3).

IC₅₀-values of compounds of the present invention are given in the following Taable.

Antithrombin activity:

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Example	IC ₅₀ (μmol·l ⁻¹)
4	0.09
24	0.01
38	0.11
40	0.02

II. Anti-factor Xa assay

Activated Factor X (Xa) is a factor in the coagulation cascade. The annti-Xa activity of compounds of the present invention was assessed by measuring spectrophotonmetrically the rate of hydrolysis of the chromogenic substrate s-2222 exterted by Xa. This assay for anti-Xa activity in a buffer system was used to assess the IC50-value of the test compound.

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In general the followed procedure and test conditions were analogous to those of the antithrombin assay as described above. Differences are indicated below.

Reference compound: benzamidine

5 Vehicle:

TNP buffer.

Solubilisation can be assisted with dimethylsulfoxide, maethanol, ethanol, acetonitrile or tert.-butyl alcohol which are without adverse effects in concentrations up to 1% (for DMSO) and 2.5% (for thee other solvents) in the final reaction mixture.

10 Technique

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Reagents*

3. S-2222 solution

One vial S-2222 (15 mg; Kabi Diagnostica, Sweden)) is dissolved in 10 ml water to give a concentration of 1.5 mg·ml⁻¹ ((2 mmol·l⁻¹).

4. Xa solution

Bovine Factor Xa Human (71 nKat·vial⁻¹; Kabi Diaggnostica) is dissolved in 10 ml TNP buffer and then further dilutted with 30 ml TNP buffer to give a concentration of 1.77 nKat·ml⁻¹. The dilution has to be freshly prepared.

Procedure

Instead of the S-2238 solution (in anti-thrombin assay), , the above S-2222 solution is added to each well in this assay.

Anti-factor Xa activity

Example	IC ₅₀ (μmol·l ⁻¹)		
1	0.64		
5	0.28		
28	0.02		

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III. Anti factor VIIa / tissue factor assay.

Vascular damage initiates a series of enzyme generation reactions ultimattely leading to the formation of a fibrin gel at the site of the injury. The primary enzyme generation reaction is the

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generation of activated factor VII (VIIa) from proenzyme factor VII. This auctivation reaction takes place by an as yet unknown mechanism. One hypothesis is that small amounts of factor Xa present in plasma, bind to the membrane-bound protein Tissue Factor (TF) -- a protein which normally does not contact blood but which gets exposed to it by injury - and 1 that this complex of membrane-bound TF and factor Xa activates factor VII (ref. 1). The activity ated Factor VII then also binds to membrane-bound TF and this intrinsic tenase complex next converts Factor X into Factor Xa.

Thrombosis develops when there is insufficient control of the coagulation reacction. One way to restore this control is by inhibiting essential coagulation enzymes such as 3 for instance the complex of membrane-bound TF and Factor VIIa. Since inhibitors of VIIaa or the VIIa/TF complex most likely will also inhibit the tenase complex, inhibitors of the latter complex may also be found by determining the inhibition of VIIa or VIIa/TF by test compounds. A method is described by which the inhibitory potency of compounds towards VIIa/TF? complex can be established. Test compounds are mixed at various concentrations with factor. VIIa and TF and with a chromogenic substrate, which is known to be split far better by TF-bound VIIa than by free VIIa. The amidolytic reaction taking place is continuously monitored in a microtiter plate reader. Inhibitory potency of the compounds investigated is expressed by that IC50, defined as the concentration of compounds yielding 50% inhibition of the amidolytice reaction, ninety minutes after the start of the reaction.

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Reagents:

Hepes buffer

A ten times concentrated Hepes buffer made by dissolving 29.40 g CaCl₂.2H₂QO, 47.66 g Hepes, 87.66 g NaCl and 30.00 g polyethyleneglycol (PEG) MW = 6000 in 1000 ml : aqua bidest. After the solution has been heated to 37 °C, the pH of the buffer is set on 7.40 withh help of 10 molar NaOH. The concentrated buffer solution is stored at 4 °C and is stable for att least two months at this condition. Prior to use the buffer is diluted in aqua bidest. 1 to 83 to obtain a final concentration in the wells (See test procedure) of 20 mM CaCl₂, 20 mM Heppes, 150 mM NaCl and 0.3 % PEG6000. If compounds are dissolved and diluted in aqua bidest. or another vehicle because of an insufficient solubility the Hepes buffer can be diluted 1 to 6 to preserve the same ionic strength in the test.

Recombinant human factor VIIa

Recombinant human factor VIIa is obtained from American Diagnostica Inc,; Greenwich, CT. Each vial contains 1.2 mg recombinant human factor VIIa, which is lyophilizedd from 2 ml buffer composed of 10 mM glycylglycine, 50 mM NaCl, 10 mM CaCl₂, 30 mg/ml1 mannitol, 0.1 % Tween, pH 5.5. The contents of each of these vials is reconstituted with 2 mml aqua bidest. as indicated by the manufacturer. The 2 ml 1.2*10⁻⁵ stock solution thus obtainined is divided in smaller fractions, which are stored at -30 °C. At this condition these VIIa sampples are stable for at least 6 months.

10 Recombinant human Tissue Factor

Recombinant human Tissue Factor is obtained from American Diagnostica Incc, Greenwich, CT. Each vial contains 25 µg recombinant human Tissue Factor (non-lipidated; MWW 35000 Dalton), which is lyophilized from 1 ml Tris/HCl buffer (pH 8.0) composed of 150 mlM NaCl, 200 mlM mannitol and 10 mlm CHAPS (Steroid derivative used to solubilize membrrane proteins; see Merck Index). The contents of each vial is reconstituted with 1 ml aqua bidesst. as indicated by the manufacturer. The 1 ml 7.14 x 10⁻⁷ M stock solution thus obtained is ddivided in smaller fractions, which are stored at -30 °C. Thus stored these VIIa samples are stable for at least 67 months.

20 Pefachrome VIIa

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Pefachrome VIIa - CH₃SO₂-D-Cha-but-Arg-pNa.AcOH (MW 670.8) - iss obtained from Pentapharm Ltd, Basle, Switzerland, in vials containing 10 µmol of this chrommogenic substrate. At the day of the experiment the contents of a vial are dissloved in 8.33 ml aquua bidest., yielding a 1.2 mMolar Pefachrome VIIa solution. What remains of this solution is storeed at -30 °C and is stable for at least 6 months at this condition.

Recombinant TF / Recombinant VIIa solution

At the day of the experiment a deep frozen sample of 1.2*10⁻⁵ M recombinannt VIIa and a deep frozen sample of recombinant human tissue factor of 7.14*10⁻⁷ is defrosted. The defrosted 7.14*10⁻⁷ solution of recombinant human TF is diluted to 4*10⁻⁷ M and 30 μH of this solution is mixed with 1 μI of the defrosted recombinant VIIa solution of 1.2*10⁻⁵ and vwith 449 μI Hepes buffer, yielding a Hepes buffer solution containing 25 nM recombinant 'VIIa and 25 nM

43

recombinant TF. The amount of 480 μl TF/VIIa solution is sufficient to examine the inhibition of eight solutions of one test compound. N times this amount is needed to estaablish the IC₅₀ of N test compounds.

5 Preparation of test compounds:

Test compounds are dissolved in Hepes buffer to give 5*10⁻³ stock solutions (A). From this solution seven additional solutions with concentrations of 1.67*10⁻³ M (B), 55.56*10⁻⁴ M (C), 1.85*10⁻⁴ M (D), 6.17*10⁻⁵ M (E), 2.06*10⁻⁵ M (F), 6.86*10⁻⁶ M (G) and 2.259*10⁻⁶ M (H) are prepared by diluting each foregoing solution with a factor three in Hepes bufferr. Such a series of solutions is prepared for the reference compound Org 34593 and also for each of the N-1 test compounds. If considered more convenient, other sets of solutions with diffferent compound concentrations may be prepared.

15 Procedure

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Compounds are distributed column by column over the microtiter plate and onee column of eight wells is reserved for a series of uninhibited reactions. Hundred μ l of Hepes buffer is brought into all (N+1)*8 wells with an eight channel pipette. Here N is the number of different test compounds, including the reference compound Org 34593. Hereafter, fifty μ l cof the pefachrome VIIa solution of 1.2 mM is added with an eight channel pipette to the 100 μ l Hepes buffer in all of the (N+1)*8 wells reserved for compound testing and the blank reactions.

Then 50 µl of each of the eight solutions of the first, second, third up to the NN-th compound is mixed in a descending order of concentrations with the contents of the first (A) until the eighth well (H) of columns 1, 2, 3, up to N respectively, so as to obtain a one compound per column distribution with a from top to bottom descending order of compound concentrations per column. Finally 50 µl Hepes buffer is added to the eight wells of the N+1 th coblumn reserved for a series of blanks.

After the whole plate has been prepared it is shaken for 1 minute in an microtiter plate shaker/incubator (Amersham) and the solutions are brought to 37 °C by incubating the plate in the same instrument for 10 minutes.

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The reactions are initiated by adding 50 µl of the 25 nM VIIa/25 nM TF scolution, which is preheated at 37 °C, to each of the (N+1)*8 wells with help of an eight channel pipette. After the plate is shaken for 30 seconds it is placed in a thermostated microtiter plate recader and the 405 nm absorbance is read in each well at time intervals of 1 minute during 90 minutes. Absorbances are collected in LOTUS 1.2.3, loaded into a PC connected to the l kinetic reader.

Evaluation:

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The (end-)absorbances measured at 90 minutes arre corrected for the blank abbsorbances at the beginning of the test by subtraction of the corresponding first absorbance vvalue measured 1 minute after the initiation of the reaction. The corrected end absorbances; in the presence (Abs[I]) and absence (Abs[O]) of the test compound are converted into) logit values by calculating +log((Abs[O]/Abs[I])-1) for each concentration [I] of the test compound. These logit values are plotted against the -log of concentrations of the test compound. Such a logit plot usually displays a lionear relationship between 20 % and 80 % inhibition of the end-absorbance.

The pIC₅₀ value is defined as the -log (concentration in M) od the test compound for which the logit value is equal to zero. This value is calculated by linear regression of thee logit vs -log [I] relation preferably around the logit zero value. When the compound tested is 550 active towards VIIa/TF that the pIC₅₀ must be calculated by extrapolation instead of interpolation, it is best to prepare an additional set of dilutions of this test compound and to perform the assay again. This method of calculating a pIC₅₀ value is described by Hafner et al. (ref. 2). The corresponding IC₅₀ is calculated as 10^{-plC50} and is expressed in Molar.

25 Quantity required:

About one mg is required to assess the IC50 of a test compound.

Reference compound:

As a reference compound Org 34593 (PPACK) may be used. For this compound an IC₅₀ of 3*10⁻⁷ M has been established.

References:

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- (1) The structural biology of expression and function of Tissue Factor: Edgingston, T.S., et al. in Thrombosis and Haemostasis 66(1), 67-79 (1991).
- (2) Mathematical analysis of concentration response relationships: Hafner, D. et al. in Arzneim.
- 5 Forsch./Drug Research 27, 1871-1873 (1977).

As a single point measurement of the anti factor VIIa / tissue factor activity of compounds of the present invention, the percentage of inhibition at a concentration of 1x10^{-5.5} M is given in the following Table. For the determination of the percentages, procedures as described above were followed.

Anti factor VIIa / tissue factor activity (percentage inhibition at a concentration of 1×10⁻⁵ M):

Example	percentage inhibbition (%)
44) EtSO ₂ -D-Phe-Leu-LysΨ[COCO]-O-	98
iPropyl	
44) EtSO ₂ -Asn-Leu-LysΨ[COCO]-O-iPropyl	56
44) EtSO ₂ -D-3-Tiq-Phe-LysΨ[COCO]-O-	91
iPropyl	
44) EtSO ₂ -D-Leu-Gln-LysΨ[COCO]-O-	94
iPropyl	

Claims:

1. A compound having the formula I

$$R^1SO_2$$
-B-X-Z-C(O)-Y (I)

5 wherein

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R¹ is R²OOC-(CHR²)_m- or R²NH-CO-(CHR²)_m- or is selected from (1-12C)alkyl, (2-12C)alkenyl, which groups may optionally be substituted with ((3-12C)cycloalkyl, (1-6C)alkoxy, OH, COOR², CF₃ or halogen, and from (6-14C)aryl, (7-15C)aralkyl and (8-16)aralkenyl, the aryl groups of which may optionally be substituted I with (1-6C)alkyl, (3-8C)cycloalkyl, (1-6C)alkoxy, OH, COOH, CF₃ or halogen;

m is 1, 2 or 3;

each group R² is independently H, (1-12C)alkyl, (3-8C)cycloalkyl,l, (6-14C)aryl or (7-15C)aralkyl, the aryl groups of which may be substituted with (1-6C)alkyl, (1-6C)alkoxy or halogen;

B is a bond, an amino-acid of the formula -NH-CH[(CH₂)_pC(O)OH]-(C(O)- or an ester derivative thereof wherein p is 1, 2 or 3, Gly, D-1-Piq, D-3-Piq, D-1-Tiqq, D-3-Tiq, D-Atc, Aic, or a L- or D-amino acid having a hydrophobic, basic or neutral side chhain;

X is an amino acid with a hydrophobic side chain, glutamine, serine, threonine, a cyclic amino acid optionally containing an additional heteroatom selected from N, O or S, and optionally substituted with (1-6C)alkyl, (1-6C)alkoxy, benzyloxy or oxo,, or X is 2-amino-isobutyric acid, -NR²-CH₂-C(O)- or the fragment

$$-NH-CH$$
 Or $N-CH_{22}-C(O)-$

wherein n is 2, 3, or 4, W is CH or N and R³ is H, (1-6C)alkyl or phenyl l which groups may optionally be substituted with hydroxy, (1-6C)alkoxy, COOH, COO(1-6CC)alkyl, CONH₂, or halogen;

Z is lysine or 4-aminocyclohexylglycine;

Y is -NH-(1-6C)alkylene- C_6H_5 , the phenyl group of which may be substituted with (1-6C)alkyl, (1-6C)alkoxy or halogen, or Y is -OR⁴ or -NR⁵R⁶, wherein RR⁴ is H, (2-6C)alkyl or benzyl, and R⁵ and R⁶ are independently H, (1-6C)alkoxy, or (1-66C)alkyl optionally

47

- 5 2. The compound of claim 1, wherein Z is lysine.
 - 3. The compound of claim 1 or 2, wherein X is a cyclic amino acid, an amino acid with a hydrophobic side chain, glutamine, serine, threonine, -NR²-CH₂-C(O)-, or thhe fragment

$$(CH_2)_n$$
-NH-CH N-CH₂-C(O)-
O
O
 $N-CH_2$ -C(O)-

wherein R³ is H, (1-6C)alkyl or phenyl.

4.. The compound of any one of claims 1-3, wherein X is proline, leucine, glutaamine, threonine, phenylalanine, -NR²-CH₂-C(O)- wherein R² is methyl, cyclopentyl or cyclohnexyl, or the fragment

-NH-CH N-CH₂-C(O)- or O
$$\mathbb{R}^3$$

wherein R³ is H or methyl.

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- 5. The compound of any one of claims 1-4, wherein B is a bond, a D-amino accid having a hydrophobic or neutral side chain.
- 6. The compound of any one of claims 1-5, wherein R¹ is (1-6C)alkyl or benzyyl.
- 7. The compound of any one of claims 1-6, wherein Y is -OCH(CH₃)₂.
- 8. A pharmaceutical composition comprising the compound of any one of claims 1-7 and pharmaceutically suitable auxiliaries.

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- 9. The compound of any one of claims 1-7 for use in therapy.
- 10. Use of the compound of any one of claims 1-7 for the manufacture of a medilicament for treating or preventing thrombin-related diseases.

INTERNATIONAL SEARCH REPORT

Inter Intel Applications No PCT/EP 98/925587

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A. CLASSIF IPC 6	CO7K5/00 CO7K5/08 A61K38/0	5
According to	International Patent Classification (IPC) or to both national classification	ion and IPC
B. FIELDS	SEARCHED	<u> </u>
IPC 6	oumentation searched (classification system followed by classification CO7K	
	ion searched other than minimum documentation to the extent that suc	
Electronic d	ata base consulted during the international search (name of data base	e and, where practical, search terms used)
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relev	vant passages F Relevant to claim No
A	JONES D M ET AL: "THROMBIN INHIB BASED ON KETONE DERIVATIVES OF AR AND LYSINE" JOURNAL OF ENZYME INHIBITION, vol. 9, 1 January 1995, pages 43-60, XP000570641 see examples 17,18; table 4 WO 96 40743 A (COR THERAPEUTICS I ;MARLOWE CHARLES K (US); SCARBORG ROBERT M) 19 December 1996	INC
	ther documents are listed in the continuation of box C.	X Patent family members are listed in annexx.
* Special c *A* docum consi *E* earlier fiting *L* docum which citati *O* docum *P* docum *P* docum	ategories of cited documents : ent defining the general state of the art which is not dered to be of particular relevance document but published on or after the international	*T later document published after the international filing date or priority date and not in conflict with the appplication but oited to understand the principle or theory ununderlying the invention. *X" document of particular relevance; the claimed 1 invention cannot be considered novel or cannot be considered to involve an inventive step when the document it taken alone of the cannot be considered to involve an invention cannot be considered to involve an inventive a step when the document is combined with one or more officer such documents, such combination being obvious to a 1 person skilled in the art. *&* document member of the same patent family
1	exclusi completion of the international search	Date of mailing of the international search repport
<u> </u>	20 July 1998	Authorized officer
Name and	Imailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswift Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,	Deffner, C-A

INTERNATIONAL SEARCH REPORT

Information on patent family members

Inter. nal Application II No PCT/EP 98/025687

NO 9640743 A 19-12-1996 AU 6590296 A 303-12-1		Information on patent family members		PCT/EP 9	PCT/EP 98/025887	
EP 0846125 A 103-06-1	Patent document cited in search report	Publication date	Patent fam member(s	nily s)	Publication c date	
	WO 9640743	A 19-12-1996	AU 659 EP 084	0296 A 6125 A	303-12-1996 103-06-1998	

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dicarbonate (425 mg) was added. The pH of the solution was adjusted and maaintained at 8 with triethylamine and the reaction was stirred for 16 hours at room temperature. Water was added and the organic layer was washed and dried to yield 782 mg of the desired product. After purification on silica using heptane/ethyl acetate 2/3 the final yield was 696 mgz.

5 TLC: Rf= 0.95, ethyl acetate/pyridine/acetic acid/water 232/31/18/7 v/v/v/v opn silica.

(b) H-Lys(Boc)Ψ[CHOHCO]-OEt.HCl

To a solution of Cbz-Lys(Boc) Y[CHOHCO]-OEt (696 mg) in ethanol (25 i mL) were added 10% palladium on activated carbon (100 mg) and 2N hydrochloric acid ((0.8 mL) and this suspension was hydrogenated at atmospheric pressure for 50 minutes at room 1 temperature. The palladium catalyst was removed by filtration and the filtrate was concentrated i in vacuo to yield H-Lys(Boc) Y[CHOHCO]-OEt. HCl (525 mg).

TLC: Rf=0.17, ethyl acetate/pyridine/acetic acid/water=232/31/18/7 v/v/v/v onn silica.

15 (c) <u>BzlSO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-OEt</u>

Coupling with BzlSO₂-norLeu(cyclo)-Gly-OH, oxidation, deprotection and purification were done according to procedures described in Example 1. Yield: 186 mg of the tittle compound. Rt (LC): 32.46 min. 20% A/80% B to 20% A/20% B/60% C in 40 min.

20 Example 4.

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BzlSO₂-norLeu(cyclo)-Gly-LysΨ[COCO]-NH₂

The coupling between BzlSO₂-norLeu(cyclo)-Gly-OH and H-Lys(Boc)\P[CH(OHCO]-NH₂.HCl. and the subsequent oxidation, deprotection and purification were done according to procedures described in Example 1 to yield 103 mg of the title compound.

25 Rt (LC): 27.50 min. 20% A/80% B to 20% A/20% B/60% C in 40 min.

Example 5.

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EthylSO₂-D-Cha-Pro-LysΨ[COCO]-OEt

The DCC/HOBt-coupling between EthylSO₂-D-Cha-Pro-OH (270 mg) and H-Lys(Boc)\P[CHOHCO]-OEt.HCl (268 mg), Dess-Martin oxidation, ddeprotection using trifluoroacetic acid and purification were done according to the procedures described in Example 1. Yield: 41 mg of the title compound.